predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

=>

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22

chain bonds :

8-11 9-12

ring bonds :

 $1-2 \quad 1-6 \quad 2-3 \quad 3-4 \quad 4-5 \quad 5-6 \quad 5-7 \quad 6-10 \quad 7-8 \quad 8-9 \quad 9-10 \quad 11-13 \quad 11-17 \quad 12-18 \quad 12-22$ 

13-14 14-15 15-16 16-17 18-19 19-20 20-21 21-22

exact bonds :

8-11 9-12

normalized bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-10 7-8 8-9 9-10 11-13 11-17 12-18 12-22

13-14 14-15 15-16 16-17 18-19 19-20 20-21 21-22

isolated ring systems :

containing 11 : 12 :

# Match level :

 $1: A \texttt{tom} \quad 2: A \texttt{tom} \quad 3: A \texttt{tom} \quad 4: A \texttt{tom} \quad 5: A \texttt{tom} \quad 6: A \texttt{tom} \quad 7: A \texttt{tom} \quad 8: A \texttt{tom} \quad 9: A \texttt{tom} \quad 10: A \texttt{tom}$ 

11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:Atom

20:Atom 21:Atom 22:Atom

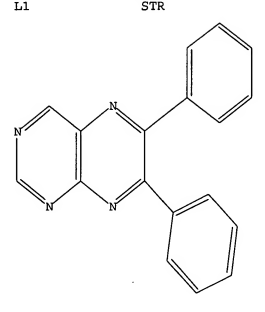
# L1 STRUCTURE UPLOADED

=> 1d

LD IS NOT A RECOGNIZED COMMAND

The previous command name entered was not recognized by the system. For a list of commands available to you in the current file, enter "HELP COMMANDS" at an arrow prompt (=>).

=> d L1 HAS NO ANSWERS L1 ST



Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 08:45:26 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 45 TO ITERATE

100.0% PROCESSED 45 ITERATIONS

16 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*
BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS:

498 TO 1302

PROJECTED ANSWERS:

80 TO 560

L2 16 SEA SSS SAM L1

=> s 11 full

FULL SEARCH INITIATED 08:45:33 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 1016 TO ITERATE

100.0% PROCESSED 1016 ITERATIONS

344 ANSWERS

SEARCH TIME: 00.00.01

L3 344 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS SINCE FILE TOTAL

ENTRY SESSION

FULL ESTIMATED COST 166.94 167.15

FILE 'CAPLUS' ENTERED AT 08:45:40 ON 30 MAY 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

Page 4 Saeed

COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 30 May 2006 VOL 144 ISS 23 FILE LAST UPDATED: 28 May 2006 (20060528/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

http://www.cas.org/infopolicy.html

=> s 13

L4 181 L3

=> file reg

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.92 168.07

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 08:47:06 ON 30 MAY 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2006 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 28 MAY 2006 HIGHEST RN 885861-83-6 DICTIONARY FILE UPDATES: 28 MAY 2006 HIGHEST RN 885861-83-6

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH January 6, 2006

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search iteration limits have been increased. See  ${\tt HELP\ SLIMITS}$  for details.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

=>

Uploading C:\Program Files\Stnexp\Queries\106792092.str

chain nodes :

24 25

ring nodes :

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22

chain bonds :

2-25 4-24 8-11 9-12

ring bonds :

1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-10 7-8 8-9 9-10 11-13 11-17 12-18 12-22

13-14 14-15 15-16 16-17 18-19 19-20 20-21 21-22

exact/norm bonds :

2-25 4-24

exact bonds :

8-11 9-12

normalized bonds :

 $1-2 \quad 1-6 \quad 2-3 \quad 3-4 \quad 4-5 \quad 5-6 \quad 5-7 \quad 6-10 \quad 7-8 \quad 8-9 \quad 9-10 \quad 11-13 \quad 11-17 \quad 12-18 \quad 12-22$ 

13-14 14-15 15-16 16-17 18-19 19-20 20-21 21-22

isolated ring systems :

containing 11 : 12 :

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:Atom

20:Atom 21:Atom 22:Atom 24:CLASS 25:CLASS

L5 STRUCTURE UPLOADED

=> d

Page 6 Saeed

L5 HAS NO ANSWERS L5 STR

Structure attributes must be viewed using STN Express query preparation.

=> s 15

SAMPLE SEARCH INITIATED 08:47:23 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 4 TO ITERATE

100.0% PROCESSED 4 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE \*\*COMPLETE\*\*

BATCH \*\*COMPLETE\*\*

PROJECTED ITERATIONS: 4 TO 200 PROJECTED ANSWERS: 0 TO 0

L6 0 SEA SSS SAM L5

=> s 15 full

FULL SEARCH INITIATED 08:47:32 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 103 TO ITERATE

100.0% PROCESSED 103 ITERATIONS 25 ANSWERS

SEARCH TIME: 00.00.01

L7 25 SEA SSS FUL L5

=> file caplus

COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST
166.94
335.01

FILE 'CAPLUS' ENTERED AT 08:47:38 ON 30 MAY 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

Page 7 Saeed

COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

Copyright of the articles to which records in this database refer is held by the publishers listed in the PUBLISHER (PB) field (available for records published or updated in Chemical Abstracts after December 26, 1996), unless otherwise indicated in the original publications. The CA Lexicon is the copyrighted intellectual property of the American Chemical Society and is provided to assist you in searching databases on STN. Any dissemination, distribution, copying, or storing of this information, without the prior written consent of CAS, is strictly prohibited.

FILE COVERS 1907 - 30 May 2006 VOL 144 ISS 23 FILE LAST UPDATED: 28 May 2006 (20060528/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

http://www.cas.org/infopolicy.html

=> s 17

L8 45 L7

=> d ibib abs hitstr tot

L8 ANSWER 1 OF 45
ACCESSION NUMBER:
DOCUMENT NUMBER:
11TLE:
11MVENTOR(5):
2005:1335074 CAPLUS
144:69859
Indoles, pteridines, pyridopyrazines, and benzotriazines as vasculostatic agents, their preparation, pharmaceutical compositions and use in therapy
Wrasidlo, Wolfgang, Doukas, John Royston, Ivor;
Noronha, Glenn Hood, John D., Dneprovskaia, Elena, Gong, Xianchang, Splittgerber, Ute; Zhao, Ningning
PATENT ASSIGNEE(5):
SOURCE:
105. Pat. Appl. Publ., 95 pp., Cont.-in-part of U.S. Ser. No. 679, 209.
CDDEN: USXXCO
DOCUMENT TYPE:

Patent English 2 DOCUMENT TYPE:

LANGUAGE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
US 2005282814	A1	20051222	US 2005-105845		20050413
US 2004167198	Al	20040826	US 2003-679209		20031002
PRIORITY APPLN. INFO.:			US 2002-415981P	P	20021003
			US 2003-440234P	P	20030114
			US 2003-443752P	P	20030129
			US 2003-463818P	P	20030417
			US 2003-466983P	P	20030430
			US 2003-479295P	P	20030617
			US 2003-679209	A2	20031002
OTHER SOURCE(S):	HARPAT	144:69859			

GΙ

$$(R^1)_m \xrightarrow{\overset{u}{\stackrel{u}{\sim}}} \overset{B}{\underset{X}{\stackrel{u}{\sim}}} (R^2)_m$$

$$\underset{H_2N}{\overset{NH_2}{\bigvee}}\underset{N}{\overset{N}{\bigvee}}\underset{N}{\overset{OH}{\bigvee}}$$

ANSWER 1 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN

#### ●2 HC1

677297-55-1P, 6,7-Bis(3-hydroxyphenyl)pteridine-2,4-diamine dihydrochloride 677297-56-2P, 6,7-Bis(3-hydroxyphenyl)pteridine-2,4-diamine methanesulfonate 677297-57-2P, 6,7-Bis(3-Hydroxyphenyl)pteridine-2,4-diamine dihydrobromide 677297-62-0P, 6,7-Bis(3,4-dihydroxyphenyl)pteridine-2,4-diamine RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(drug candidate; preparation of vasculostatic agents and use for

treatment of disorders associated with compromised vasculostasis)

RN 677297-55-1 CAPLUS

CN Phenol, 3,3'-(2,4-dismino-6,7-pteridinediyl)bis-, dihydrochloride (SCI)
(CA INDEX NAME)

677297-56-2 CAPLUS Phenol. 3,3'-(2,4-diamino-6,7-pteridinediyl)bis-, methanesulfonate (salt) (SCI) (CA INNEX NAME)

CRN 677297-51-7 CHF C18 H14 N6 O2

Page 9 Saeed

ANSWER 1 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)
The invention relates to nitrogen heterocyclic compds. of formula I, which
are useful for treating disorders associated with compromised vasculostasis.
In compds. I, each of A, B, V, X, Y, and Z is independently selected from
C, C(0), N, and NR3, where R3 is H or (un)substituted alkyl, each R1 is
independently halo, OR4, N(R4)2, or SR4, where R4 is H, lower alkyl, aryl,
heteroaryl, etc.; each R2 is independently halo, OR5, N(R5)2, SR5, OPO3H2,
(un)substituted alkyl, (un)substituted aryl, (un)substituted heteroaryl,
where R5 is H, lower alkyl, aryl, heteroaryl, etc.; and each of m and n is
independently an integer from 1 to 4. The invention also relates to the
preparation of I, pharmaceutical compns. comprising a compound I and a
pharmaceutically acceptable carrier, as well as to the use of the compns.
for the treatment of a variety of disorders including stroke, myocardial
infarction, cancer, ischemis/reperfusion injury, autoinmume diseases such
as rheumatoid arthritis, eye diseases such as retinopathies or macular
degeneration, inflammatory diseases, vascular leakage syndroms, edens,
transplant rejection, adult/acute respiratory distress syndroms (ARDS),
and the like. Cyclocondensation of 3,3"-dihydroxybenzil with
2,4,5,6-tetraaminopyrimidine sulfate results in the formation of
diaminopteridine II. Compound II expresses an ICSO value of 83 nM in an
assay for the inhibition of the human pl20y subunit of P13 kinase
and results in 65% reduction of myocardial infarction in rats.
677237-51-7P, 6,7-Bis(3-hydroxyphenyl)pteridine-2,4-diamine
dihydrochloride
RL: PAC (Pharmacological activity), RCT (Reactant), SSN (Synthetic
preparation), TRU (Therapeutic use), BlOL (Biological study), PRRP
(Preparation), PACT (Reactant or reagent), USES (Uses)
(drug candidate) preparation of vasculostatic agents and use for

treatment

iment
 of disorders associated with compromised vasculostasis)
677297-51-7 CAPLUS
Phenol, 3,3'-(2,4-diamino-6,7-pteridinediyl)bis- (9CI) (CA INDEX NAME)

677297-63-1 CAPLUS 1,2-Benzenediol, 4,4'-(2,4-diamino-6,7-pteridinediyl)bis-, dihydrochloride (SCI) (CA INDEX NAME)

ANSWER 1 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

677297-57-3 CAPLUS Phenol, 3,3°-(2,4-diamino-6,7-pteridinediyl)bis-, dihydrobromide (9CI) (CA INDEX NAME)

677297-62-0 CAPLUS
1,2-Benzenediol, 4,4'-{2,4-diamino-6,7-pteridinediyl}bis- (9CI) (CA INDEX

677298-35-0, 6,7-Bis-(3-hydroxyphenyl)pteridine-2,4-diamine

677298-35-0, 6,7-Eis-(3-hydroxyphenyl)pteridine-2,4-diamine
sulfate
RL: PAC (Pharmacological activity), THU (Therapeutic use), BIOL
(Biological study), USES (Uses)
 (preparation of vasculostatic agents and use for treatment of disorders
associated with compromised vasculostasis)
677298-35-0 CAPLUS
Phenol, 3,3'-(2,4-diamino-6,7-pteridinediyl)bis-, sulfate (salt) (9CI)
(CA INDEX NAME)

CH 1

CRN 677297-51-7 CMF C18 H14 N6 O2

CH. 2

CRN 7664-93-9 CMF H2 O4 S

LØ ANSWER 2 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:308364 CAPLUS DOCUMENT NUMBER: 140:321386

INVENTOR (5):

140:321366
Preparation of vasculostatic agents and methods of use Vrasidlo, Volfgang, Doukas, John; Royston, Ivor; Noronha, Glenn; Hood, John D., Dnsprovskaia, Elena; Gong, Xianchang, Splittgerber, Ute; Zhao, Ningning Targegen, Inc. USA
PCT Int. Appl., 230 pp.
CODEN: PIXXU2
Patent
English
2

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.					KIND DATE				APPL	ICAT								
WO 2004030635				A2 20040415				WO 2	003-		20031002							
WO 2004030635																		
	V:									BB	BG,	22	BY	B7	C	C)	CN1	
			CR.	CII.	CZ.	DR.	DK.	DM.	DZ.	EC.	EE,	EG.	ES.	FI.	GR.	GD.	GE.	
											KE.							
											MN,							
											SE,							
											VN.						ın,	
	DIT.	GH,															24	
	W.																	
											CH,							
											NL,							
											G₩,							
	2500				AA		2004	0415		CA 2	003-	2500	727		2	0031	002	
					A1 20040423													
EP	1549																	
	R:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	5K		
BR	2003	0150	53		λ		2005	0809		BR 2	003-	1505	3		2	0031	002	
CN	1720 2006	224			Α		2006	0111		CN 2	003-	8010	4711		2	0031	002	
JP	2006	5153	17		T2		2006	0525		JP 2	005-	5003	78		2	0031	002	
DRITE	( APP	LN.	INFO	. :						US 2	002-	4159	81P		P 2	0021	003	
										us 2	003-	4402	34P		P 2	0030	114	
											003-					0030		
										us 2	003-	4638	182		2	0030		
											003-							
											003-					0030		
											003-					0031		
PD 61	MIDO	/e1 .			MADI		140.	2212		2	003-				- 2	5031	UU2	

OTHER SOURCE(S): MARPAT 140:321386

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

Compds. (2 Markush structures shown as I and II; others are described in the claims and disclosure; variables defined below; e.g. III and IV) and methods are provided for treating disorders associated with compromised vasculostasts. Invention methods and compon, are useful for treating a variety of disorders including for example, stroke, myocardial infarction, cancer, ischemia/reperfusion injury, autoimnume diseases such as retinopathies or nacular degeneration or other vitreoretinal diseases, inflammatory diseases,

Page 10 Saeed

L8 ANSWER 1 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN

ANSWER 2 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) vascular leakage syndrome, edema, transplant rejection, adult/acute respiratory distress syndrome (ARDS), and the like. Although the methods of prepn. are not claimed, many example prepns are included. For example, III was prepd. (75 %) from 2-(2-aminophenyl) incle and 4-hydroxyphenylacetic acid. Various expts. are described that show the use of the claimed compds. along with chemotherapeutic agents for cancer treatment. The claimed compds. along with chemotherapeutic agents for cancer treatment. The claimed compds. along who inhibition of vascular leak induced by interleukin 2. Inhibition of VEGF-induced edema, redn. of myocardial infarction and inhibition of c-Src and Yes kinases were demonstrated for some of the claimed compds. For I: each RO = -H, -COOH, -OR', -SOHM, wherein R' is -H or lower alkyl, or when x = 2, each RO is taken together to form a 1,3-dioxolyl ring, or each RO = (un) substituted alkyl, (un) substituted alkyl, (un) substituted arylalkyl, or -4-hydroxyphenyl. 677297-51-78, 6,7-Bis (3-hydroxyphenyl) pteridine-2,4-diamine 677297-63-18, 6,7-Bis (3,4-dihydroxyphenyl) pteridine-2,4-diamine 677237-63-1P, 6,7-Bis (3,4-dihydroxyphenyl)pteridine-2,4-diamine dihydrochloride
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or respect); USES (USes) (drug candidate; preparation of vasculostatic agents and methods of use) 677297-51-7 CAPLUS
Phenol, 3,3'-(2,4-diamino-6,7-pteridinediyl)bis- (9CI) (CA INDEX NAME) L8 ANSWER 2 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

RN 677297-63-1 CAPLUS
CN 1,2-Benzenediol, 4,4'-{2,4-diamino-6,7-pteridinediyl}bis-, dihydrochloride
(9CI) (CA INDEX NAME)

●2 HC1

IT 18181-93-69, 6,7-Diphenylpteridine-2,4-diamine
677297-50-69, 6,7-Bis(3-hydroxyphenyl)pteridine-2,4-diamine
monohydrochloride 677297-55-19, 6,7-Bis(3hydroxyphenyl)pteridine-2,4-diamine dihydrochloride 677297-56-29
, 6,7-Bis(3-hydroxyphenyl)pteridine-2,4-diamine methanesulfonate
677297-51-39, 6,7-Bis(3-hydroxyphenyl)pteridine-2,4-diamine
dihydrobromide 677297-62-09, 6,7-Bis(3,4dihydroxyphenyl)pteridine-2,4-diamine 677298-35-09,
6,7-Bis(3-hydroxyphenyl)pteridine-2,4-diamine sulfate
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PRPE (Preparation); USES
(Uses)
(drug candidate; preparation of vasculostatic agents and methods o

(Uses) (drug candidate; preparation of vasculostatic agents and methods of use)
RN 18181-93-6 CAPLUS
CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

L8 ANSWER 2 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) (9CI) (CA INDEX NAME)

OH 1

CRN 677297-51-7 CMF C18 H14 N6 O2

CH 2

CRN 75-75-2 CMF C H4 03 S

RN 677297-57-3 CAPLUS
CN Phenol, 3,3'-(2,4-diamino-6,7-pteridinediyl)bis-, dihydrobromide (9CI) (CA INDEX NAME)

●2 HBr

RN 677297-62-0 CAPLUS CN 1,2-Benzenedio1, 4,4'-(2,4-diamino-6,7-pteridinediy1)bis- (9CI) (CA INDEX Page 11 Saeed

L8 ANSWER 2 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continue

RN 677297-50-6 CAPLUS
CN Phenol, 3,3'-(2,4-diamino-6,7-pteridinediyl)bis-, monohydrochloride (9C1)
(CA INDEX NAME)

• HCl

RN 677297-55-1 CAPLUS CN Phenol, 3.3'-(2,4-diamino-6,7-pteridinediyl)bis-, dihydrochloride (9CI) (CA INDEX NAME)

●2 HC1

RN 677297-56-2 CAPLUS CN Phenol, 3,3'-(2,4-diamino-6,7-pteridinediyl)bis-, methanesulfonate (salt)

L8 ANSWER 2 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) NAME)

RN 677298-35-0 CAPLUS
CN Phenol, 3,3'-(2,4-dismino-6,7-pteridinediyl)bis-, sulfate (salt) (9CI) (CA INDEX NAME)

CM

CRN 677297-51-7 CMF C18 H14 N6 O2

CM.

CRN 7664-93-9

•

но-9-0

L8 ANSWER 3 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2003:758726 CAPLUS DOCUMENT NUMBER: 140:314320 Determination of lipophilic de

2003:788726 CAPLUS
140:314320
140:314320
1a0:314320
1a0

AUTHOR (S):

SOURCE:

Blomedical Chromatography (2003), 7(16), 365-372

CODEN: BICHEZ: ISSN: 0269-3879

FUBLISHER: John Wiley & Sons Ltd.

JOURNAI TYPE: Journal

LANGUAGE:

AB The liquid chromatog, retention factors extrapolated to pure water, k'w, for several 6,7-diaryl-pteridine derivs. in both an octadecylsilane (ODS) and an immobilized artificial membrane column (IMM.PC.DD2), using action trile-aqueous buffer pH = 7.45 as mobile phase, were obtained. The logarithms of the k'w values in the IAM.PC.DD2 column, log k'IAMw, show good correlation with the calculated values of the octanol-water partition coeffs., log Po/w, showing that the chromatog, parameter can be used as lipophilicity descriptor for the studied pteridines. However, interactions other than the lipophilic ones seem to be involved in the ODS column. Previous studies have shown that pteridines have antibelminic properties. In spite of the complexity of the studied biol. system as compared with the chromatog, one, good correlation between the descriptors obtained in the IAM column and biol. activity (expressed as the log of the inhibitory concentration required to obtain up to 50% in the reduction of population population growth of nematodes, log ICSO) was observed IT 18181-93-6

CORPORATE SOURCE:

IT 18181-93-6
RI: ANT (Analyte); BSU (Biological study, unclassified); PAC (Pharmacological activity); PRP (Properties); ANST (Analytical study); BIOL (Biological study); (determination of lipophilic descriptors of antihelmintic 6,7-diaryl-pteridine derivs. for bioactivity predictions)
RN 18181-93-6 CAPUS
CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

18181-93-6 CAPLUS 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

L8 ANSWER 4 OF 45 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 2003:125984 CAPLUS DOCUMENT NUMBER: 138:368664

DOCUMENT NUMBER:

ACCESSION NUMBER: 2003125994 CAPLUS
DOCUMENT NUMBER: 138:368664
TITLE: Pteridines. Part CXIII. Protection of pteridines
AUTHOR(S): Fachbereich Chemie, Universitaet Konstanz, Konstanz,
D-78457, Germany
SOURCE: Helvetica Chimica Acta (2003), 86(1), 1-12
CODEN: HACACAV: ISSN: 0018-018
DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASERACT 138:36864
AB The low solubility of pterins can drastically be improved by N2-acylation or formation of the N2-[(dimethylamino)methylene] derivs. Both types of compds, can be albylated under Mitsunobu conditions to form N2-acylpterins and their derivs. N2, N2-Dimethylpterins and N2-methylpterins direct alkylation to the 04-position. Deacylation can be achieved under very mild conditions by solvolysis with MeGR, and displacement of the O4-position.

the corresponding pteridin-2,4-dismines. Cleavage of the N2[(dimethylamico)methylene) group works well with ammonia. The advantage of applying the 2-(4-nitrophenyl) ethyl (npe) group as blocking group is seen in its selective removal by 1.8-diazabicyclo[5.4.0]undec7-ene (DBU) under aprotic conditions without harming the other substituents.

RE: SPN (Synthetic preparation), PREF (Preparation) (N2-acylation of pterins followed by Mitsunobu alkylation to form alkyl derivs. of pterins)

1818-19-3-6 CAPLUS

2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 5 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN

(Continued)

REFERENCE COUNT:

60 THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 6 OF 45
ACCESSION NUMBER:
DOCUMENT NUMBER:
TITLE:
APPLICATION
AUTHOR(S):

AUTHOR(S):
CAPIUS COPYRIGHT 2006 ACS on STN
1998:237746 CAPIUS
128:291449
Application of neural networks to the study of structure-activity relationships of 6,7-disrylpteridines as nematocides
Ochos, C., Rodriguez, J., Rodriguez, H., Chana, A.,
Stud, M., Alonso-Villalobos, P., Martinez-Grueiro, M.

CORPORATE SOURCE:

Inst. Quimica Medica, Madrid, 28006, Spain
Stud, M.; Alonso-Villalobos, P.; Martinez-Grueiro, M.

Inst. Quimica Medica, Madrid, 28006, Spain
SOURCE:

Medicinal Chemistry Research (1997), 7(9), 530-545
CODEN: MCREED; 155N: 1054-2523

PUBLISHER:
Birkhaeuser Boston
DOCUMENT TYPE:
LANGUAGE:
AB A study of structure-activity relationships of 6,7-diarylpteridines as nematocides, using a trained back-propagation neural network, has been carried out. This network has sallowed the prediction of the qual.
nematocide activity of pteridine derivs. not yet synthesized. Among 25 preselected pteridine derivs. 17 were predicted as nematocides by the network. The synthesis and the nematocidal activity of the pteridines, which had been predicted as active compds., are reported. Use of this network allows the prediction of qual. nematocide activity of pteridine derivs., not yet synthesized.

Itials-3-6

RL: AGR (Agricultural use); BIOL (Riologica)

REFERENCE COUNT:

19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 8 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
1995:324866 CAPLUS
1171LE:
1171LE

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

	PAT	ENT	NO.			KIN	D	DATE			APP	LICAT	ION	NO.		D	ATE		
							-									-			
	80	9427	439			λl		1994	1208		WO	1994-	U544	74		1	9940	425	
		V:	AT,	AU,	BB,	BG,	BR,	BY,	CA,	CH,	CN	, CZ,	DE,	DK,	ES,	FI,	GB,	GE,	
			HU,	JP,	KG,	KP,	KR,	KZ.	LK,	LU,	LV	, MD,	MG.	MN,	MW,	NL,	NO,	NZ,	
			PL,	PT,	RO,	RU,	SD,	SE,	SK,	TJ,	TT	, UA,	UZ,	VN					
		RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IE,	IT,	LU,	MC,	NL,	PT,	SE,	
			BF.	BJ,	CF,	CG,	CI,	CH,	GA,	GN,	ML	, MR,	NE,	SN,	TD,	TG			
		5521				Α		1996	0528		US	1993-	6789	7		1	9930	527	
	λU	9467	726			A1		1994	1220		AU	1994-	6772	6		1	9940	425	
	US	5532	367			Α		1996	0702		US	1995-	4160	17		1	9950	331	
	US	5639	753			A		1997	0617		US	1995-	6126	57		1	9951	128	
RIO	RIT	Y APP	LN.	INFO	.:						US	1993-	6789	7		A 1	9930	527	
											WO	1994-	U544	74	1	7 1	9940	425	
THE	R SC	DURCE	(S):			MARI	PAT	122:	2586	55									

Pteridine and 8-deazapteridine compds. and compns. were prepared and used for controlling insects in agricultural crops. These pteridines may be represented by structure [I, R and RI = NHZ, lower alkylamino, dl(lower alkylamino, el.,g., NH2), or di(lower alkylaminotechyleneanine (e.g., N=CHH42); RZ = H, NHZ, lower alkyl (e.g., -CH3, -CH(CH3)2), di(lower alkyl) anionaethyleneanino, CH, lower alkyn, Ph or substituted Ph, haloslkylphenylalkyl (e.g., 3-trifluoronaethylphenylnethyl); Q = N or CH; R3 = (n)=R4, n = 0 or 1; when n = 1, n is a bridging stom or noiety selected from O, S, SO, SOZ, lower alkylene (e.g., CHZ or CHZCH2), lower alkynlene (e.g., CHZ-CH3), clover alkynlene (e.g., CHZ-CH3), or clover alkynlene (e.g., CHZ-CH3), or clover alkynlene (e.g., CHZ-CH3), or clover haloslkenylene (e.g., CHZ-CH3), CO, anionaethyl (e.g., CHZ-CH3), or (substituted amino)methyl (e.g., CHZ-CH3), and R4 = H, lower alkyl (e.g., He, i-Pr), thien-2-yl, pyridin-3-yl, or II; V, Y, X, Y = H, halo, haloslkyl, aryl, Ph, PhO; Z = H or halo]. A typical dust formulation against tohacco budworm contained 1 part 2,4-diamino-6-[3,5-di(trifluoronaethyl)phenyl]-7-nethylpteridine and 99 parts talc.

Page 13 Saeed

L8 ANSWER 7 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
1996:466654 CAPLUS
125:157774
Anthelaintic activity of 6,7-diarylpteridines
AUTHOR(\$):
CORPORATE SOURCE:

synthesized from the corresponding diaminopyrimidines and account aldehydes.

Their anthelmintic activity was tested in vitro against Caenorhabditis elegans and Heligmosomoides polygyrus and in vivo against Trichinella spiralis. Structure-activity relationships are discussed.

IT 18181-93-6F
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SFN (Synthetic preparation); BIOL (Biological study); PREF (Preparation)

(anthelminic activity and preparation of disrylpteridines)

RN 18181-93-6 CAPLUS
CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

ANSWER 8 OF 45 CAPLUS COPYRIGHT 2006 ACS on SIN (Continued)
10101-93-6P
RL: AGR (Agricultural use); SPN (Synthetic preparation); BIOL (Biological
study); PREP (Preparation); USES (Uses)
(preparation and insecticidal activity of pteridine derivs.)
18101-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

L8 ANSWER 9 OF 45 CAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER: 1994:134419 CAPLUS
DOCUMENT NUMBER: 120:134419
TITLE: Protection

120:134419
Protection and deprotection of fused
2-maino-4(3H)-pyrimidinones: conversion of pterins and
5-deazapterins to 2,4-diamino derivatives
Taylor, Edvard C., Otty, S. R., Durucasu, Inci
Dep. Chem., Princeton Univ., Princeton, NJ, 08544, USA
Hetercoycles (1993), 36(8), 1883-95
CODEM: HICYAM; ISSN: 0385-5414

AUTHOR (5): CORPORATE SOURCE:

DOCUMENT TYPE:

English CASREACT 120:134419 LANGUAGE: OTHER SOURCE(S):

AB 5-Dearapterins and pterins are readily converted to their 4-deoxy-4-amino derivs., e.g. I, (a lactam-to-amidine conversion) by reaction with 4-chlorophenyl phosphorodichloridate and 1,2,4-triazole to give intermediate 4-[1'-(1,2,4-triazoly1)] derivs., e.g. II, followed by reaction with aqueous ammonia. Some anomalous results obtained by application of the Mitsunobu reaction (normally a lactam-to-lactim ether conversion) to 5-dearapterins are detailed.

II 18181-93-69

18181-93-69
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)
18181-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

ANSWER 10 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Co 18181-93-6 CAPLUS 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME) (Continued)

151648-52-1 CAPLUS 2,4-Pteridinediamine, 6,7-bis(4-aminophenyl)- (9CI) (CA INDEX NAME)

$$\underset{H_2N}{\overset{NH_2}{\longrightarrow}}\underset{N}{\overset{NH_2}{\longrightarrow}}\underset{NH_2}{\overset{NH_2}{\longrightarrow}}$$

L8 ANSWER 10 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1994:4377 CAPLUS
DOCUMENT NUMBER: 120:4377 CAPLUS
11TLE: 120:4377 CAPLUS

DOCUMENT TYPE: LANGUAGE: AB Toxoplasma

CODEN: AMACCUP 133N: VOUNT-05-13

NAME: Journal UAGE: Regish
TOXOPlasma gondi: RH was obtained in high yield from culture in RPMI medium on a line of Chinese hamster ovary cells lacking dihydrofolate reductase activity (ATCC 3952 dhfr-). Dihydrofolate reductase prepns. from harvested organisms had pp. activities of 22.9 mol/kain/ag. The 50% inhibitory concms. against reference compds. were 0.014 pM for Orevatas.

inhibitory concan. against reference compds. were 0.014 µM for methotrexate,

0.25 µM for pyrimethamine, 2.7 µM for trimethoprim, and 0.010 µM for trimetrexate. The Km value for NADPH was 11 µM and followed Michaelis-Menten kinetics; the Km for dihydrofolate was .apprx.11 µM, but substrate inhibition appeared to occur at high substrate concas. Dihydrofolate reductase from T. gondi was used to screen 130 compds. from the National Cancer Institute repository. Thirteen compds. were >100-fold more potent than pyrimethamine toward T. gondi dihydrofolate reductase; 6 compds. with various potencies were 8-46 times as selective as pyrimethamine for the protocoal form of the enzyme over the mammalian form. Four trimetrexate analogs were more potent than trimetrexate, and 2 were significantly more selective. Representative compds. were also tested in a culture model of T. gondi employing uracil incorporation as an index of growth. One pyrimethamine analog was as effective as pyrimethamic in inhibiting T. gondi in culture (50% inhibitory concentration,

0.45 µM). Three other compds. were also effective at micromolar concns.

6967-77-7 18181-93-6 151648-32-1

6967-77-7 18181-93-6 131648-52-1
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study) (dihydrofolate reductase of Toxoplasma gondii inhibition by, structure in relation to) 6967-77-7 CAPLUS
Phenol, 4,4'-(2,4-diamino-6,7-pteridinediyl)bis- (9CI) (CA INDEX NAME)

L8 ANSWER 11 OF 45
ACCESSION NUMBER:
DOCUMENT NUMBER:
1976:164728 CAPLUS
1976:164728 CAPLUS
84:164728
Direct conversion of 4-hydroxypteridines to their
4-amino analogs
Gapaki, G. R., Whiteley, J. M.
SCHIPPS Clin. Res. Found., La Jolla, CA, USA
Chem. Biol. Pteridines, Proc. Int. Symp., 5th (1975),
627-32. Editor(s): Pfleiderer, Wolfgang. de Gruyter:
Berlin, Ger.
CODEN: 32LMAC
Conference

Conference

DOCUMENT TYPE: LANGUAGE: GI English

The 4-aminopteridines I (R = NH2, Rl = H, Me, Ph) were prepared in 27-428 yields by treating I (R = OH) with PhOP(o) (NH2) 2. isla1-93-69
RL: SFN (Synthetic preparation); PREP (Preparation) (preparation by amination of hydroxypteridines with phenylphosphorodiamidate) 18181-93-6 CAPLUS 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

ΙT

L8 ANSWER 12 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1974:37071 CAPLUS
80:37071
TITLE: Pteridines. LVIII. Synthesis and properties of pterin and 2,4-diaminopteridine mono- and di-N-oxides
Yamamoto, Hiroshi; Hutzenlaub, Volfgang; Pfleiderer,
Volfgang
CORPORATE SOURCE: Pachbereich Chem., Univ. Konstanz, Constance, Fed.
Rep. Ger.
COEN: CHEEAM; ISSN: 0009-2940

DOCUMENT TYPE: Journal
German
GI For diagram(s), see printed CA Issue.
AB Treating pterins (i, n = m = 0) R, R1 = H, Me, or Fh) and
diaminopteridines (II, n = m = 0) with H202-CF3CO2H gave preferentially
the oxides I and II (n = 1, n = 0, and n = m = 1), the structures of which
were proven by rearrangement, hydrolysis, and uv spectra: I (s = n = 0, R
= H, R1 = CMe3 and Fh) were oxidized to give the 5-oxides I (n = 1, n = 0)
AUS COENTIAL OF THE PROPOSED OF THE PROPO

51324-31-3 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl-, 5,8-dioxide (9CI) (CA INDEX NAME)

L8 ANSWER 14 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1968:458202 CAPLUS
OCCURENT NUMBER: 25:58202 CAPLUS
69:58202
Stimulation by pteridines of the uptake of anethopterin by human lymphocytes
AUTHOR(S): Kessel, Davids Botterill, Viviennes Hall, Thomas C.
CORFORATE SOURCE: Lab. of Pharmacol., Children's Cancer Res. Found.,
BOSTON, HA, USA
BOCOMENT TYPE: JOURNAL STAN. 60:6-2952
DOCUMENT TYPE: JOURNAL STAN. 60:6-2952
LANGUAGE: English

LANGUAGE:

MAGNI TYPE: Journal
RUAGE: English
Triamterene (2,4,7-triamino-6-phenylpteridine) and certain other
pteridines stimulated the uptake of amethopterin by human small
lymphocytes, apparently by removing a barrier to amethopterin transport.
This stimulation did not extend appreciably to other cell types or to
other lymphocyte transport systems tested. 19 references.
BLE NEW STATES.

18101-93-6
RL: BIOL (Biological study)
{
amethopterin absorption by lymphocytes in response to)
18191-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (SCI) (CA INDEX NAME)

L8 ANSWER 13 OF 45 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1969:446130 CAPLUS DOCUMENT NUMBER: 71:46130

DOCUMENT NUMBER

71:46130
Dihydrofolate reductase from Trypanosoma equiperdum.
II. Inhibition by 2,4-diaminopyrimidines an related heterocycles
HcCormack, John J., Jr., Jaffe, Julian J.
Coll. of Hed., Univ. of Vermont, Burlington, YT, USA Journal of Medicinal Chemistry (1969), 12, 662-8
CODEN: JMCHAR, ISSN: 0022-2623
Journal
Emplish TITLE:

AUTHOR(S): CORPORATE SOURCE: SOURCE:

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE: Journal
LANGUAGE: English
AB A number of 2,4-diaminopyrimidines and related heterocyclic compds. have
been

evaluated as inhibitors of dihydrofolate reductase obtained from T. equiperdum, chicken liver, and rat liver. 2,4-Diaminopyrimidine itself (at 10-4M) was not effective as an inhibitor of dihydrofolate reduction in

(at 10-4M) was not effective as an inhibitor of dihydrofolate reduction in 3 systems studied but 5-aryl derive. of 2,4-diaminopyrimidine were good in-inhibitors (IDSO - 10-8 to 10-6M) of this enzymic reaction.
2,4-biamino-5-benzylpyrimidines and 2,4-diamino-5-aryloxypyrimidines were considerably more effective as inhibitors of the trypanosomal enzyme system than of the mammalian and evian systems. Although 2,4-diamino-6-phenyl-s-triazine was not active as an inhibitor of the enzymes studied, related 4,6-diamino-1,2-diphenylpteridine was found to be approx. 10-fold more effective as an inhibitor of the 3 reductase systems than was 2,4-diamino-6,7-dimethylpteridines 2-amino-6,7-diphenylpteridine and 4-amino-6,7-diphenylpteridines 2-amino-6,7-diphenylpteridines and 4-amino-6,7-diphenylpteridines ever not effective as inhibitors of these enzymes. 2,4,7-Triamino-6-arylpteridines bearing an ortho substituent in the 6-aryl molety were 10-100-fold more potent as inhibitors of the reductase systems than were the corresponding para-substituted derivs. The 2-amino-4-hydroxypteridine derivs. biopterin, xanthopterio, and isoxanthopterin were effective neither as substrates nor as inhibitors of the trypanosomal reductase.

RL BIOL (Biological study)
(tetrahydrofolate dehydrogenase response to)

18181-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

L8 ANSWER 15 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1968:452104 CAPLUS
DOCUMENT NUMBER: 69:52104

AUTHOR(5): Pteridine diuretics
Weinstock, Joseph, Vilson, James V., Wiebelhaus,
Virgil D., Nams, Alfred R., Brennan, Francis T.,
Sosnowski, Genevieve
CORPORATE SOURCE: Res. and Develop. Div., Smith Kline and French Lab.,
Philadelphis, PA, USA
Journal of Medicinal Chemistry (1968), 11(3), 573-9

COUNTENT TYPE: JOURNAM, ISSN: 0022-2623
JOURNAM, ISSN: 0022-2623
JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM, ISSN: 0022-2623

JOURNAM,

INITIATION REPORT REPOR

L8 ANSUER 16 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1960:118339 CAPLUS
DOCUMENT NUMBER: 54:118339
RIGINAL REFERENCE NO.: 54:22664-1, 22665a-b
Fittle: Pteridines. XXI. One-step synthesis of
4-aninopteridines
AUTHOR(S): Taylor, Edward C., Jr., Cheng, C. C.
CORPORATE SOURCE: Princeton Univ., Princeton, NJ
Journal of Organic Chemistry (1959), 24, 997-9
CODEN: JOCEAN: ISSN: 0022-3263
DOCUMENT TYPE: Journal of Organic Chemistry (1959), 24, 997-9
CODEN: JOCEAN: ISSN: 0022-3263
DOCUMENT TYPE: Unavailable
CASPRACT 54:118339
GI For diagram(s), see printed CA Issue.
AB cf. CA 54, 5675s. The title compds., N: CR.N:C(NH2).C:C.N:CR'.CR':N(I),
were prepared by heating amidine selts of (NC) 2C:NOH (II) in HOCH2CH2OH,
diluting the isomerized pyrimidine solution with H2O, adding Na25204.2H2O
(III)

and finally treating with an o-diketone. II K salt (1.0 g.) and 1.1 g. guanidine carbonate (IV) in 10 ml. HOCHZCHZOH warmed gently 3 mln. as the deep red solution diluted with 10 ml. HZO, 0.6 g. III added and the

mixture

heated 20 min. on a steam bath, the clear yellow solution acidified to pH 6

with HCl and warmed 15 min. on a steam bath with 1 ml. Ac2, diluted with 20

ml. alc., and the chilled solution filtered yielded 75% authentic I (R =

NH2, R' = Me) (V). II K salt (1.5 g.), 1.5 g. IV, and 12 ml. HOCHZCH2OH heated 3 min. and reduced with 1.0 g. III, the alkaline solution refluxed 1 hr. with 10 ml. EtCOMe in 5 ml. alc., and the filtered solution chilled yielded 29% I (R = NH2, R' = Ph) (VI). II K salt (3 g.) and 3.3 g. IV in 20 ml. hot HOCHZCH2OH reduced with 1.6 g. III and the pale yellow solution adjusted to pH 3 with Hcl, stirred 40 min. at 110 with 7.5 g. glyoxal bisulfite in 50 ml. H2O and the mixture kept overnight at room temperature, acidified with AcOH, and the separated product (3.55 g.) sublimed at 240 % 0.05 mm. gave authentic I (R = NH2, R' = H) (VII). II K salt (2.0 g.) and 2.2 g. IV isomerized and reduced, the solution diluted with 20 ml.

NH HCl and treated with 2.0 g. alloxan, the purple mixture shaken with gradual separation of an orange solid and adjusted to pH 9 with KOH, the alkaline

gradual separation of an orange solid and adjusted to pH 9 with KOH, the line solution heated 10 min. at 110° and reacidified to pH 6 with HCl, refrigerated, and filtered gave 76% orange 2,4-diamino-5,7-dibydroxypyrimido[5,4-g]pteridine (VIII), m. above 350°. II benzamidine salt (2.0 g.) and 10 ml. 2-picoline heated 30 min. at 135° and the mixture diluted with 20 ml. H2O, the blue-green suspension evaporated in vacuo and the residue heated to 90-100° in 25 ml. H2O, treated portionwise with 1.6 g. III and the yellow solution stirred 20 min., refluxed 2 hrs. with 2 g. Bz2 in 30 ml. 1:1 ECCOMe-alc., and the cooled mixture filtered gave 1.9 g. 4-amino-2,6,7-triphenylpteridine (IX), m. 255°, & 290, 377 mg (log e 4.53, 4.23, alc.).
Paper chromatographic analysis in a series of systems by the descending method at 22° gave fluorescent spots (pteridine, Rf with 44 Na citrate, 3% NH4Cl, 2:1 SN BuCH-AcOH, and 2:1 PrOH-1% NH4OH given): V, 0.25, 0.54, 0.43, 0.55; VI, 0.08, 0.17, 0.73, 0.88; VII, 0.27, 0.54, 0.28, 0.51; VIII, 0.23, 0, 0, 0; IX, 0, -, 0.88, 1. Except as indicated, I

L8 ANSWER 17 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1959:40178 CAPLUS
ORIGINAL REFERENCE NO.: 53:40178 CAPLUS
ORIGINAL REFERENCE NO.: 53:72611,7262a-b
ITITLE: Effect of 4-amino folic antagonists on biological acetylations
AUTHOR(S): Johnson, Willard J., Corte, George, Jasmin, Roland
CORPORATE SOURCE: F. W. Horner Labs., Montreal, Can.
Proceedings of the Society for Experimental Biology and Medicine (1958), 99, 677-80
CODEN: PSERBAJ ISSN: 0037-9727
DOCUMENT TYPE: Journal
LANGUAGE: Unavailable
AB cf. C.A. 50, 15662f. Acetylation of sulfanilamide and of isoniazid in pigeon-liver exts. was markedly inhibited (noncompetitively) by 4-amino analogs of folic acid. Amethopterin (10-5M) and aminopterin (5 + 10-5M) inhibited acetylation about 60H. Folic acid (10-3M) was not inhibitory, and failed to reverse the inhibition by Amethopterin.
2,4-Diamino-6,7-diphenylpteridine (10-3M) gave 51 inhibition and 2,4-diaminopteridine (10-3M) was inactive. Amethopterin, administered to rabbits cojointly with sulfanilamide, resulted in a marked increase in plasma level of free sulfanilamide with concomitant decrease in acetylaufianilamide. The possible significance of these results with regard to combination chamotherapy of cancer is discussed.

IT 18101-93-6, Pteridine, 2,4-diamino-6,7-diphenyl(acetylation inhibition in liver by)

RN 18181-93-6 CAPLUS
CN 2,4-Pteridineddamine, 6,7-diphenyl(SC) 2,4-Pteridineddamine, 6,7-diphenyl(SC) 2,4-Pteridineddamine, 6,7-diphenyl(SC) 2,4-Pteridineddamine, 6,7-diphenyl(SC) 2,4-Pteridineddamine, 6,7-diphenyl-

ANSWER 16 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) prepd. by this one-step method were chromatographically pure. 1818-19-3-6, Pteridine, 2,4-diamino-6,7-diphenyl-(preparation of) 18181-9-3-6 CAPLUS 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

IΤ

L8 ANSWER 18 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1956:52652 CAPLUS
DOCUMENT NUMBER: 50:52652
RIGINAL REFERENCE NO.: 50:101039=9
ITILE: Route to 4-aminopteridines
AUTHOR(S): Taylor, E. C., Jr., Paudler, W. W.
CORPORATE SOURCE: Princeton Univ., Princeton, NJ
Chemistry & Industry (London, United Kingdom) (1955)
1061-2
CODEN: CHINAG, ISSN: 0009-3068
DOCUMENT TYPE: Journal
LANGUAGE: Unavailable
CASREACT 50:52652
AB A new route for 4-amino-5,6-diphenylpyrazines (III) is described.
2-Bydroxy-5,6-diphenylpyrazinamide (III) (Jones, C.A. 43, 3009h) gave 998
yield 2-chloro-3-cyano-5,6-diphenylpyrazine (III) when heated in a sealed
tube with PC13. III was also obtained in 80% yield by heating a mixture of
II, POC13, and PC15. Pusion of III with guanidine carbonate, urea, or
thiourea gave 65, 59, and 51% 2-amino, 2-hydroxy, and 2-mercapto derivaof I, resp. III with NZH4.HZO gave 2-chloro-5,6-diphenylpyrazinediphenylpyrazinemide when treated with NXH0H and XI, or
2-amino-3-cyano-5,6-diphenyl-1-pyrazolo[b]pyrazine. III gave 2-amino-5,6diphenylpyrazinamide when treated with NXH0H and XI, or
2-amino-3-cyano-5,6-diphenylpyrazine when fused with NXHOAC.
II 18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl(Preparation of)
RN 18181-93-6 CAPLUS
CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

ANSWER 22 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)
Phenol, 4,4°-(2,4-diamino-6,7-pteridinediyl)bis- (9CI) (CA INDEX NAME)

18181-93-6 CAPLUS
2,4-Pteridinedismine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

151648-52-1 CAPLUS
2,4-Pteridinediamine, 6,7-bis(4-aminophenyl) - (9CI) (CA INDEX NAME)

804555-05-3 CAPLUS Acetanliide, 4'-[7-(p-acetamidophenyl)-2,4-diamino-6-pteridyl]- (5CI) (CA INDEX NAME)

L8 ANSWER 22 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN

857398-11-9 CAPLUS Pteridine, 2,4-diamino-6,7-bis[m-nitrophenyl]- (5CI) (CA INDEX NAME)

L8 ANSWER 22 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

855404-13-6 CAPLUS INDEX NAME NOT YET ASSIGNED

857397-78-5 CAPLUS
Pteridine, 2,4-diamino-7-(p-nitrophenyl)-6-phenyl- (SCI) (CA INDEX NAME)

857397-80-9 CAPLUS Pteridine, 2,4-diamino-6-(p-nitrophenyl)-7-phenyl- (5CI) (CA INDEX NAME)

857398-09-5 CAPLUS
Pteridine, 2,4-diamino-6,7-bis[p-nitrophenyl]- (5CI) (CA INDEX NAME)

L8 ANSWER 23 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1955:17275 CAPLUS 49:17275
DOCUMENT NUMBER: 49:17275
A9:17275 CAPLUS 49:17275
A9:17275 CAPLUS 49:17275
A9:17275 CAPLUS 50:17275
A9:17275 CAPLUS 50:1727

L8 ANSWER 19 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1955:84700 CAPLUS
ORIGINAL REFERENCE NO.: 49:160141,16018-b
Synthesis of compounds related to thymine. II. Effect
of thymine antagonists on the biosynthesis of DNA
(deoxyribonucleic acid)
AUTHOR(5): Bardos, Thomas J., Levin, Georgia M.; Herr, Ross R.;
Gordon, Harry L.
CORPORATE SOURCE: Armour Labs., Chicago
Journal of the American Chemical Society (1955), 77,
4279-86
CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Unavailable
Lactobacillus leichmannii and L. plantarum. The modes of action of
various metabolic antagonists, particularly 5-bromouracil and its
nucleosides, are discussed. The systems described are used to study the
biol. action of 3 new thymine antagonists: 5-arcaptouracil, 5-uracilyl
disulfide, and uracil-5-isothiouronium chloride. Deoxyuridine (2.28 g.)
in 50 cc. water treated with saturated Br water, the solution serated,
lyophilized, the residue in 250 cc. absolute EtOH refluxed 15 min., and
concentrated
in vacuo to 30 cc. yielded 0.82 g. 5-bromodeoxyuridine, m. 181-3\*.

IT 18181-93-6 CAPLUS
CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

L8 ANSWER 21 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1955:29379 CAPLUS
OCCUMENT NUMBER: 49:29379
ORIGINAL REFERENCE NO.: 49:56781,5679a-b
ACTION ACCESSION COMMENT
TITLE: Action of 2.4-disamino-6.7-diisopropylpteridine upon
Plasmodium gallinaceum and its relation to other
compounds which are pteroylglutamic acid antagonists
Bishop, Ann
Univ. Cambridge, UK
Parasitology (1954), 44, 450-64
CODEN: PARARS; ISSN: 0031-1820
JOURNET TYPE: Journal
LANGUAGE: Unavailable
AB Two strains of P. gallinaceum were made resistant to 2,4-diamino-6,7diisopropylpteridine (1) by passing the parasite through chicks which had
been treated with the drug. Passages at 2-4-day intervals maintained a
state of acute infection, and large nos. of the parasites were exposed to
the drug. Dosages were kept slightly below the maximum tolerated by the
parasite. Strains resistant to I were resistant to proguani (II),
pyrimethanine (III), 2,4-diamino-5-(7-diphenylpteridine; but not to
sulfadiazine (V). In one strain, development of resistance to II was
developed at a faster rate than resistance to II, and resistance to II was
resistant to I, II, and III, but not to V. The action of I and II was not
antagonized by p-aminobenoic acid, though in the min. effective dose
their action was antagonized by relatively large doses of pteroylglutamic
acid but only by pteroylglutamic acid when the V was given in small doses.

11 1618-193-6 (Parlid on 2,4-diamino-6,7-diphenyl(effect on Plasmodium gallinaceum)

N 1818-193-6 (APJUS

CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

L8 ANSWER 20 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1955:44060 CAPLUS
ORIGINAL REFERENCE NO.: 49:8484a-c
Pittle: Phenolic compounds as chemotherapeutic agents against poliosyselitis virus in tissue culture
AUTHOR(5): Kramer, Patricia Elly, Robbins, Mary Louise, Smith, Paul K.
CORPORATE SOURCE: George Washington Univ., Washington, DC, USA
Journal of Pharmacology and Experimental Therapeutics (1955), 113, 262-71
CODEN: JETAB, ISSN: 0022-3565
JOURNAT TYPE: Journal Unavailable
AB Tests were made on 135 phenolic compds. and 20 nonphenolic benzene derivs.
Only 19 compds, were found to inhibit proliferation of type 2
poliomyelitis virus, Y-SK strain, in roller cultures of monkey testicular tissue inoculated at the same time with drug and virus. Fifteen (all diphenols or aminophenols) were capable of inhibiting virus-induced degeneration of the fibroblasts over a wide range of concentration, independent

degeneration of the fibroblasts over a wide range of concentration independent of whether the virus was inoculated 24 h. before or after treatment with the drug. Twenty-six compds. naturally occurring in tissues were tested for ability to reverse the inhibitory action of the drugs. Glutathione reversed the action of 12. Sarine, threonine, and hydroxyproline frequently inhibited the action of one or more of the drugs.

IT 6867-77-7, Pteridine, 2,4-diamino-6,7-bis[p-hydroxyphenyl][effect on poliomyelitis virus in tissue culture]

RN 6967-77-7 CAPLUS

CN Phenol, 4,4'-(2,4-diamino-6,7-pteridinedlyl)bis- (9CI) (CA INDEX NAME)

L8 ANSWER 22 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1955:20270 CAPLUS
ORIGINAL REFERENCE NO.: 49:40307-1,4031a
Z1TLE: 2,4-Diaminopteridine and derivatives
INVENTOR(s): Cornelius K.
PATENT ASSIGNEE(s): DECUMENT 179E: Unavailable
PANHLY ACC. NUM. COUNT:
PATENT INFORMATION: 1

CAPLUS COPYRIGHT 2006 ACS on STN
1955:20270 CAPLUS
CAPLU

PATENT NO. KIND DATE APPLICATION NO. DATE

US 2667486 19540126 US 1951-228139 19510524
A new series of antibacterial compds., 2,4-diaminopteridine, also called 2,4-diaminopyrimido(4,5-b)pyrazine, and its substitution products, are prepared from 2,4,5,6-tetrasminopyrimidine (I) or its saits with 1,2-dicarboxyl compds., 1,2-dicarboxylic acids, or with a-carbonyl acids or their derivatives such as esters, etc., in aqueous, nonaq., or d

of solns. of acidic, neutral, or basic reaction. On the pyrimidine ring are located 2 amino groups, at the 2- and the 4-position, making this new synthetic pterin to be the first to have a 2,4-diamino structure, unlike folic acid. Thus, 2,4-diaminopteridine was prepared by adding 2 g. I sulfate in 70 cc. hot H2O, to 3.5 g. of glyowal bisulfite in 30 cc. of hot water, bolling the clear yellow mixture is bmin. treating with C, allowing to cool slowly, filtering off the light yellow microcryst. precipitate, inc

Vacer, Bolling the Clear yellow mixture is min., treating with C. including to cool slowly, filtering off the light yellow microcryst. precipitate, hing with water and He2CO, drying in vacuo, and purifying by recrystn. from H2O or by sublination at 180'/1 mm. Other compds. prepared are 2,4-diamino-7-methylpteridine; 2,4-diamino-7-methylpteridine; 2,4-diamino-7-methylpteridine; 2,4-diamino-7-methylpteridine; 2,4-diamino-7-methylpteridine; 2,4-diamino-6,7-diphylorycypteridine; 2,4-diamino-6,8-dipylorycypteridine; 2,4-diamino-6,8-dipylorydinylyrimido[4,5-b]pyrazine; 2,4-diamino-6,7-diphylorycypteridine; 2,4-diamino-6,7-diphylorycypteridine; 2,4-diamino-6,7-diphylorycypteridine; 2,4-diamino-6,7-diphylorycypteridine; 2,4-diamino-6,7-diphylyrimido[4,5-b]pyrazine; 2,4-diamino-6,7-diphylyrimido[4,5-b]pyrazine; 2,4-diamino-6,7-diphylyrimido[4,5-b]pyrazine; 2,4-diamino-6,7-diphylyrimido[4,5-b]pyrazine; 2,4-diamino-6,7-diphylyrimido[4,5-b]pyrazine; 2,4-diamino-6,7-diphylyrimido[4,5-diphylyrimido]pyrazine; 2,4-diamino-6,7-diphylyrimido[4,5-diphylyrimido[4,5-diphylyrimido]pyrazine; 2,4-diamino-6,7-diphylyrimido[4,5-diphylyr

(preparation of) 6967-77-7 CAPLUS

L8 ANSWER 24 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1954:3619 CAPLUS
ORIGINAL REFERENCE No.: 48:3619
ORIGINA DOCUMENT TYPE:

LANGUAGE:

LOUDEN: JACSAT, ISSN: 0002-7863

DOCUMENT TYPE:

LANGUAGE:

L due acidified with AcOH yielded 1.09 g. 4-amino-2-mercapto-6,7-diphenylpteridine (VI), m. 283°. III and benzil yielded 474 4-amino-2-methylmercapto-6,7-diphenylpteridine (VII), m. 252.5-53°. VI (0.15 g.) in 50 cc. absolute ECOH and 0.10 g. MeI refluxed 30 min., the solution evaporated nearly to dryness in vacuo, the residue in 40 cc. 0.1N v evaporated to dryness, the residue in 50 cc. boiling EtOH treated with C and the filtrate diluted with 50 cc. water yielded 0.110 g. VII, m. 252.5-53. Approx. 0.20 g. pteridine, 1.0 g. maine, and 50 cc. absolute EtOH heated 10 hrs. at 180° yielded the following alkylpteridines (pteridine, amine, product, % yield, and m.p. given): IV, NH3, I, 78, 200-3° (uncor.): V, NH3, I, 79, -V, VI, MeNHZ, 4-amino-2-methylamino-6,7-diphenylpteridine (VIII), 97, 264-5°; VII, MeNHZ, VIII, 68, -V VI, MeNHZ, 4-amino-2-dimethylamino-6,7-diphenylpteridine (IX), 93, decompose above 260°; V, MeNHZ, X, 53, -. VI (1.0 g.), 15 cc. piperidine, L8 ANSWER 25 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1952:60851 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 46:60851 CAPLUS CORGINAL REFERENCE NO.: 46:10212d-g Improvements in the control of t FATENT ASSIGNEE(S):
DOCUMENT TYPE:
PARENT ASSIGNMENT (S):
DOCUMENT TYPE:
PATENT INFORMATION:

46:10212d-g
Improvements in the preparation of sulfonyl derivatives of piperazine
Societe des usines chimiques de Rhone-Poulenc
Parent INFORMATION:

46:10212d-g
Improvements in the preparation of sulfonyl derivatives of piperazine
Societe des usines chimiques de Rhone-Poulenc
Unavailable
1

PARENT INFORMATION: APPLICATION NO. KIND DATE GB 661537 19511121 GB

For diagram(s), see printed CA Issue.

Piperazine derivs. of the general formula RN(CH2.CH2) 2NSO2 (CH2) nSO2N(CH2.C H2) ZNSO2 (CH2) nso2N (CH2.C H2) ZNSO2 (CH2) ZNSO2 (CH2.C H2) ZNSO2 (CH hrs. at room temperature, the precipitate centrifuged and washed with Et20; hrs. at room temperature, the precipitate centrifuged and washed with Et20 the I (n = 5, R = Me) so obtained, purified by extraction with Et20 in a Soxhlet apparatus and recrystn. from Et0Ac, m. 102-3°. The following compds. I were prepared in a similar manner (n, R, and m.p. given): 3, Et, 97-98°, 4, Et, 171°, 5 Et, 110-11°, 6, Et, 14°. The disulfonyl halides [II] may be prepared by treating Br(CH2)nBr with NH2CSNH2, which gives the thiocarbamido derivative, (NBr.NN:)NH2CS(CH2)nCSC(NH2):NH.HBr (III). This latter product the KOAC gives the corresponding discetate [IV]. II is formed by treating IV in water with a halogen. The III so prepared are (n and m.p. given): 5, 170°, 3, 205°, 4, 215°, 6, 205°. IV: 3, 46°, 4, 29-4°, 6, 86°. These compds. are effective in the treatment of states of traumatic or hemorrhagic shock.

11 18181-93-6, Pteridine, 2,4-diamino-6,7-diphemyl(preparation of)

ANSWER 24 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) and 10 cc. HCOMMe2 refluxed 12 hrs., the soln. distd. in vacuo, the residue poured into 100 cc. water, and the oil in a few cc. He2CO poured into 50 cc. ice water yielded 0.75 g. 4-amino-2-piperidino-6,7-diphenylpteridine m. 209°. VI (0.70 g.) and 10 cc. morpholine refluxed 10 hrs., the mixt dild. with water, let stand overnight at 2°, filtered, the product in Me2CO treated with C and the filtrate evapd. to drynes yielded 0.37 g. the 2-morpholino analog, m. 231-2°.
18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl-(alkyl derivs.)
18181-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME) (prepn. of

L9 ANSWER 26 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1952:60850 CAPLUS DOCUMENT NUMBER: 46:60850 ORIGINAL REFERENCE NO.: 46:10212d Pyrimidopyrazines Timmis, Geoffrey M. Wellcome Foundation Ltd. INVENTOR (S): INVENTOR(S): T:
PATENT ASSIGNEE(S): P:
DOCUMENT TYPE: P:
LANGUAGE: U:
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION: Unavailable PATENT NO. KIND DATE APPLICATION NO. GB 674847 19520702 GB 5ee U.S. 2,581,889 (C.A. 46, 7594g).
18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl(preparation of)
18181-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

18181-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

L8 ANSVER 27 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1952:57451 CAPLUS
ORIGINAL REFERENCE NO.: 46:57451
ORIGINAL REFERENCE NO.: 46:5823d-i
TITLES: 6- and 7-Bromomethyl pteridines
BOOTHE, James H.
ABFICAN CAPLUS
LANGUAGE: Unaveilable
PATENT INFORMATION:

APPLICATION NO. KIND DATE US 2584538 19520205 US 1948-35069 19480624 Compds. Useful as intermediates in the preparation of pteroylglutamic acid

and related compds. are prepared as 2-Amino-4-hydroxy-6-methylpteridine [II] 12 g., 48% HBr 400, and Br 12 cc. are heated overnight on a steam bath, the mixture chilled overnight, and the crystals isolated mechanically and crystallized from 48% HBr by a current of air-Br, and then from hot AcOH

crystallized from 48% HBF by a CUTrent of air-Bc, and then from hot ACON taining
1-5% HBF, to give 2-amino-4-hydroxy-6-(dibromomethyl)pteridine-HBF (III).
II 50 g., Br 50, and 48% HBF 1500 cc. are heated 20 hrs. on a steam bath, and the solution concentrated to 500 cc., chilled at -5° overnight, filtered, concentrated to 550 cc., and cooled to give 38 g. III. The final filtrate is evaporated to dryness, and the residue suspended in 1 l. H20, shaken, collected, and vashed with H20, alc., and Et20 to give 6-(bromomethyl)pteridine, which with N-(p-aminobenzoyl)glutamic acid gives 9-27% I. II 20 g., 48% HBF I l., and Br 12 cc. are refluxed 1 hr., the solution concentrated to 500 cc., treated with 21 g. C. filtered, added to

cold H2O, and neutralized to pH 3-5 with NaOAc to give 24 g. III (free base). Br 50 g. in 48% HBr 300 cc. is added dropwise with stirring to 2-amino-4-hydroxy-7-methylpteridine (IV) 50 g. in 48% HBr 3 l., and the mixture heated 20 min. to give 2-amino-4-hydroxy-7-(bromomethyl)-pteridine. Br 40 cc. is added to IV 40 g. in 48% HBr 1200 cc. at 70-95\*, and the solution heated 2 hrs. on the steam bath to give 47.7 g. 2-amino-4-hydroxy-7-(dibromomethyl)-pteridine (V). IV 2 g., 48% HBr 40 cc., and Br 4 g. are heated 45 min. just under reflux temperature, the tion solution

freed of excess Br, cooled, and the precipitate suspended in H2O containing

ral drops of pyridine to give 2.3 g. V. Br 1.34 cc. in 48% HBr 10 cc. added to 2-amino-4-hydroxy-6,7-dimethylpteridine 5 g. at 95° gives 3.5 g. 2-amino-4-hydroxy-6-methyl-7-(bromomethyl)-pteridine. (BrCH2CO)2 1.22 g. in alc. 10 cc. added to 2.4,5-triamino-6-hydroxypyrimidine-2HCl in 2.5 N HBr 25 cc. gives 0.9 g. of a substance of the same composition as 6,7-dibromodinethylpterin (VI). H2O is added to VI 15 g. in 48% HBr 210 cc. to give 750 cc. of solution KI 7.13 g. in H2O 25 cc. is added during 1 hr. at 55°, and the solution held 1 hr. at 55°, cooled to 15°, treated with NaHSO3 until the dark color is discharged, filtered, and neutralized to pH 1 with saturated NaOAc solution to give g.

12.1 g.

2-amino-4-hydroxy-6-bromomethyl-7-methylpteridine. Br 250 mg. is added to
2-amino-4-hydroxy-6-methylpteridine 200 mg. in (HOCH2) 2 7.5 cc. and 48%

L8 ANSWER 28 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1952:57450 CAPLUS
OCCUMENT NUMBER: 46:57450
GORIGINAL REFFERENCE NO: 46:9623b-d
SUBSTITUTE: Substituted pteridines
Campbell, Norman R.; Fitzgerald, Maurice E. H.;
COLLIER, Henry O. J.
Allen & Hanburys Ltd.
PATENT ASSIGNEE(S): DOCUMENT TYPE: Unavailable
FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

GB 656769 19510829 GB
AB Compds. which have antibacterial properties, especially against Vibrio cholerae, are prepared as follows. A mixture of (EtCO)2 1,2,4,5,6-tetraaminopyrimidine acetate (1) 1 mol., and 60% AcOH 2.5.1 are refluxed 2 hrs., cooled, poured into 20 1. HZO, and adjusted to pH 6 to give 6,7-diethyl-2,4-diaminoptoridine (II), mc 20° (from EtOH). Similarly prepared are the following analogs of II: 6,7-di-iso-Pr, m. 246°s anixture of 7,6- and 6,7-Et(p-MacCGHs), mc 228°s amixture of 7,6- and 6,7-Et(p-MacCGHs), mc 220°s 6,7-di-Pr, m. 200°s 6,7-di-Pr, m. 200°s 6,7-di-Sec-Bu, mc 210°s a mixture of 7,6- and 6,7-iso-Pr, hn. 242°s amixture of 1.12503 4, anisil 5 g., HaCORt 40, HZO 80, EtOH 40, and HCl 2.4 cc. is refluxed 10 hrs., filtered, neutralized with NaOR solution, and cooled

ed
to give the 6,7-(p-MeOC6H4)2 analog of II, m. 281° (from pyridine).
The same compound prepared similarly to II m. 288°.
694514-86-8, Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)(preparation of)
694514-86-9 CAPLUS
Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)- (5CI) (CA INDEX NAME)

ANSWER 27 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)
HBF 0.5 cc. at 50°, the soln. heated 45 min. at 50-70°,
mixed with (HOCH2) 2 7.5 cc. contg. N-(p-aminobenzoyl) glutamic acid 0.5 g.,
buffered at pH 4 with 1 g. KOAc, and heated overnight at 100° to
give 0.2 g. of naterial contg. 8.23% I. Cf. C.A. 46, 3032d.
694514-86-8, Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)(preparation of)
694514-68-8 CAPLUS
Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)- (SCI) (CA INDEX NAME) ΙT

L8 ANSWER 29 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1952:45488 CAPLUS
CORIGINAL REFREENCE NO.: 46:45488
G16145489 CAPLUS
46:45488 CAPLUS
46:

PATENT NO. PATENT NO. KIND DATE APPLICATION NO. DATE
US 2581889 19520108 US 1949-103319 194901
Nitrosomminopyrimidines in acidic media with a ketone containing an 19490706

AB Nitrosominopyrimidines in actors means are unequivocally activated -CH2- group yield pyrimidopyrazines whose structures are unequivocally known. E.g., 2 g. 5-nitroso-2,4,6-triaminopyrimidine, 4 g. PhcH2COPh (I), 60 cc. glacial RDAc, and 1 drop concentrated RCI are heated 9 hrs. at 150-60°, cooled, the yellow solid filtered, the filtrate diluted with 250 cc. H20 and 20 cc. 2 N HCl, shaken twice with 50 cc. Et20, then with 50 cc. light petr. ether to remove unchanged I, the solution made alkaline with

50 cc. light petr. ether to remove unchanged 1, the solution made alkaline with concentrated NH4OH, and the precipitate filtered, washed with H2O and MeOH, and dried,
yielding 1.1 g. 2.4-diamino-6.7-diphenylpyrimido-[4,5-b] pyrazine (C.A. numbering throughout), m. 822' (from 508 HCOZH). Similarly prepared are 2.4-diamino-6H-indolo[2,3-g]pteridine, does not m. under 350';
2.4-diaminopyrimido[4,5-b] pyrazine, m. 332'; 2.4-diamino-6-phenyl-7-methylpyrimido[4,5-b] pyrazine, m. 332'; 2.4-diamino-6-methyl-7-phenylpyrimido[4,5-b] pyrazine, m. 332'; 2.4-diamino-6-methyl-7-phenylpyrimido[4,5-b] pyrazine, m. 332'; 6.8-di-aminodipyrimido[4,5-b,5,5',4'-e] pyrazine-2,4-6.010 dipyrimido-[4,5-b,5',4'-e] pyrazine-2,4,6.010 dipyrimido-[4,5-b,5',4'-e] pyrazine-2,4,6.8(1M,3H,7H,9H)-tetrone, m. 403'. These compds. are useful therapeutic agents.

IT 18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl(preparation of)
NN 18181-93-6 CAPLUS
CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

La ANSWER 30 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1952:36322 CAPLUS
DOCUMENT NUMBER: 46:36322
ORIGINAL REFERENCE NO: 46:6197-ad
TITLE: The activities of some 2,4-diaminopteridines and sulfathiazole against Streptococcus faecalis and Staphylococcus aureus
AUTHOR(S): Collier, H. O. J. J. Waterhouse, Pamela D.
AUTHOR(S): Allen & Hamburys Ltd., Ware, UK
British Journal of Pharmacology and Chemotherapy (1952), 7, 161-9
COUENT TYPE: Journal
LANGUAGE: Unavailable
AB cf. C.A. 45, 5815i. In vitro tests were made with 6,7-disubstituted 2,4-disaminopteridines as growth inhibitors against 4 strains of S. faecalis. The greatest activity was shown by dislkyl derivs. with straight or branched chains containing 3-6 C, the disheryel (I) derivative, and

straight or branched chains containing 3-6 C, the dibenzyl (1) derivative,

1'-methyl-indolo-(2',3',6,7)-2,4-diaminopteridine. Highest activity was
shown sainst strains requiring preformed pteroylglutamic acid (II).

Sulfathiazole (III) potentiated the inhibitory effect of I against strains
of S. faecalis not requiring II. The presence of 58 urine or oxelated
horse blood did not appreciably antagonize the inhibitory effect of I.

Against S. aureus, I. bis(cyclohexylatehyl), and normal dialkyl compds.

were most active. In the dialkyl series peak activity occurred in the
dibutyl and diamyl derivs. The toxicity of I was similar to that of III.
I phosphate prolonged the lives of mice infected with S. aureus. It also
acted synergistically with III both in vitro and in vivo in protecting
mice against infections of S. aureus.

18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl6851248-68-6, Pteridine, 2,4-diamino-6,7-bis[o-methoxyphenyl]887228-94-5, Pteridine, 2,4-diamino-6,7-bis[o-methoxyphenyl](antibacterial action of)

18181-93-6 CAPLUS

2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

694514-86-8 CAPLUS
Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)- (5CI) (CA INDEX NAME)

L8 ANSWER 31 OF 45 CAPLUS COPYRIGHT 2006 ACS ON STN ACCESSION NUMBER: 1952:27482 CAPLUS DOCUMENT NUMBER: 46:27482
ORIGINAL REFERENCE NO.: 46:4675b-e

46:4675b-e
2, 4-Diaminopyrimidines. A new series of antimalarials
Falco, E. A.; Goodwin, L. G.; Hitchings, G. H.; Rollo,
I. M.; Russell, P. B.
Wellcome Research Labs., Tuckahos, NY
British Journal of Pharmacology and Chemotherapy
(1951), 6, 185-200
CODEN: BJPCAL; ISSN: 0366-0826 TITLE: AUTHOR (S):

CORPORATE SOURCE:

DOCUMENT TYPE: LANGUAGE:

(1951), 6, 185-200
CODEN: BAPCAL: ISSN: 0366-0826
JUENT TYPE: Journal
JUAGE: Unavailable
of. Nature 164, 1133 (1949). Many members of a series of 158 derivs. of
cf. Nature 164, 1133 (1949). Many members of a series of 158 derivs. of
cf. Nature 164, 1133 (1949). Many members of plasmodium gallinaceum in
chicks and Plasmodium berghei in mice. Compds. with a 5-Ph substituent
were most active. 5-PhcR2 and 5-Pho derivs. were somewhat less active.
Substitution of a p-No2 or p-halogen in the 5-substituent enhanced the
activity. Substitution of an elkyl group in the 6-position enhanced the
activity and in the 5-Ph derivs. maximum activity was reached with the 6-Et
compound 2, 4-Dianino-5-(p-chlorophenyl)-6-ethylpyrindine was 60 times as
active as paludrine against P. gallinaceum and 200 times as active against
P. berghei Longer-chain alkyl derivs. were less active. The drugs were
active against the blood forms of Plasmodium cynomolgi in monkeys, but had
no pronounced action on the excepthrocytic forms. Acute oral LD50
(mg./kg.) in mice, and the paludrine equivs. against P. gallinaceum and P.
berghei were, in order, for the following more active 2,
4-diaminopyrimidines: 5-(p-chlorophenoxy)-6-methyl. apprx. 1000, 0.4, 0.7;
5-(p-chlorobenzyl)-6-methyl. 79, 0.4, 2.0, 5-(p-chlorophenyl), 250, 0.4,
30; 5-(p-nitrobenzyl)-6-methyl, 19, 0.4, 2.0; 5-(p-chlorophenyl), 250, 0.4,
30; 5-(p-nitrobenzyl)-6-methyl, 19, 0.4, 2.0; 5-(p-chlorophenyl)-6-ethyl,
22, 60, 200; and 5-(3, 4-dichlorophenyl)-6-ethyl, 66, 20, 190, while
quinine gave 1160, 0.08, 0.16, and paludrine acetate 59, 1.0, 1.0.
500284-43-5 CAPLUS
2,4-Pteridinediamine, 6,7-bis(4-chlorophenyl)- (9CI) (CA INDEX NAME)

L8 ANSWER 30 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN

857228-94-5 CAPLUS
Pteridine, 2,4-diamino-6,7-bis(o-methoxyphenyl)- (5C1) (CA INDEX NAME)

L8 ANSWER 32 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1952:17637 CAPLUS
DOCUMENT NUMBER: 46:17637
ORIGINAL REFERENCE NO: 66:3059g-1,3060a-h
AUTHOR(S): ANALOGS of Pteroylglutamic acid. VII. 2-Alkylamino
derivatives
AUTHOR(S): Roth, Barbara; Smith, James M., Jr.; Hultquist, Martin

AUTHOR(S):

Roth, Barbara's Smith, James M., Jr.; Hultquist, Martin E.

CORPORATE SOURCE:

An. Cyanamid Co., Bound Brook, NJ
Journal of the American Chemical Society (1951), 73, 2864-8

CODEN: JACSAT, ISSN: 0002-7863

DOUMENT TYPE:
LANGUAGE:

AB cf. C.A. 44, 9451e.

NAME (22.7 q.) in 60 cc. MeOH added to 25.8 g.

MeZNC(:NH)NHZ in 50 cc. dry MeOH, 20.7 g. NCCH2COZMe added to the refluxing mixture during 10 min., and the mixture refluxed 3 hrs., filtered, and neutralized with HCl yielded 26.0 g. 2-dimethylanino-4-hydroxy-6-aminopyrimidine (1), m. 290.5-25. (from water). I (2 g.) in 20 cc. water warmed and acidified, the pH adjusted to 4 with NaOAc, then 0.74 g. NaNOZ in 2 cc. water added slowly at 80', yielded the 5-nitroso derivative (11) of 1, n. 259' (decomposition). NaSZO4 (10 g.) (ITA) added at 50' to 5 g. II in 30 cc. water containing a min. of dilute NaOH yielded 3.5 g. 2-dimethylamino-4-hydroxy-5,6-diaminopyrimidine sulfite (111). III (20 g.) in 330 cc. water acidified with HCl, then warmed in vacuo, 10.7 g. N. (p-aminobenzoyl)qlutamic acid (17) added, the pH adjusted to 3.0 with NaOH, 3.98 g. NaZCC207 in 23 cc. water and 17.3 g. CHZPECHBCHO (V) in 16 cc. AcCH added dropvise and simultaneously during 20 min. to the mixture at 45' (pH maintained at 3), and the mixture after 20 min. at 45' cooled to 10' yielded 21.3 g. assaying 24.88 N-(p-[(2-dimethylamino-4-hydroxy-6-pteridylmethyl)amino] benzoyl-glutamic acid (VI). Purification by solution and precipitation yielded VI, assaying 85.18. Ac2 (1 g.) and 2.9 g. III in 30 cc. water heated 45 min.

led VI,
assaying 85.18. Ac2 (1 g.) and 2.9 g. III in 30 cc. water heated 45 min.
at 85', cooled, and neutralized with NH4OH yielded
2-dimethylamino-4-hydroxy-6,7-dimethylpteridine, m. 283-8' (from
alc.) (decomposition). NH2C(:NH)NHCN (VIA) (50 g.) and 100 g. MeZNH,HCl

2-dimethylamino-4-nyurusy-0, "Name why special College (decomposition). MHCC(INN) NHCN (VIA) (50 g.) and 100 g. HeZNH.HCl heated

3 hrs. at 180°, the mixture poured into 600 cc. absolute EtOH, the solution cooled to 10°, filtered, 58.5 g. NaOHe added, then 66.7 g. CH2(CN)2 dropwise during 20 min. to the refluxing mixture, and the mixture refluxed 2 hrs., cooled, filtered, and the product washed with ice water yielded 61 g. 2-dimethylamino-4, 6-diaminopyrindidine (VII), m. 259-60° (from dilute alc.). H2504 (5 N) added to 10 g. VII in 200 cc. water to obtain solution, the pH adjusted to about 4 with NaOAc, and 25 NaNO2 added to the solution at 85° to a permanent starch-KI test yielded 11.2 g. 2-dimethylamino-4, 6-diamino-5-nitrosopyrindidine (VIII). VIII (42.9 g.) in 550 cc. water adjusted to pH 2.5 with 5 N HCl, 130 g. IIA added slowly at 60°, and the mixture heated to 70°, then acidified to approx. pH 2 with dilute H2504, yielded 56 g. 2-dimethylamino-4, 5,6-triaminopyrindidine sulfate (IN). IX (21.3 g.) and 19.5 g. BaCl2.2H2O in 330 cc. water warmed 10 min. to 60°, cooled to 45°, then treated with IV and V as for VI yielded N-(p-(2-dimethylamino-4-amino-6-pteridylnethyl) amino)benzoyl)glutamic acid (X). VIII (5 g.) and 4.59 g. BaCl2 in 50 cc. water heated on the steam bath 10 min., filtered hot, and 1.62 g. Ac2 added yielded 2-dimethylamino-4-amino-6-7-dimethylpteridina-HCl, bright yellow crystair from dilute alc. MeHHZ.HCl (14) and 620 g. VIA heated 3 hrs. at 180°, the melt cooled to 100°, poured into 6 1. absolute EtOH, 1,440 g. NacNew added to the filtrate, then 1,280 g. NCCH2CO2Et during 30 min., and the mixture refluxed 4 hrs. yielded 550 g.

ANSWER 32 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)
3-mathyl-2,6-diamino-4(3M)-pyrimidone (XI), m. 265-72' (depending on the rate of heating): the cooled filtrate yielded 20 g. addnl. material: the filtrate cond. to 2 l. yielded 442 g. 2-mathylamino-4-hydroxy-6-aminopyrimidine (XII), m. 227-9' (from sic. and water).
2-Mathylmsrcapto-4-hydroxy-6-aminopyrimidine (Johns and Baumann, C.A. 7, 2288) (10 g.) and 40 cc. 25 MeHRE heated 5 hrs. at 120' yielded e white cryst. product, m. 245-7.5' nitroso deriv., CGH9N502. XII (9 g.) nitrosated yielded 9.5, orange ppt. (XIII) which did not m. helow 360'. XIII (9 g.) reduced with 22 g. IIA yielded (XIV), CGH9N502. XII (9 g.) nitrosated yielded 9.5, g. orange ppt. (XIII) which did not m. helow 360'. XIII (9 g.) reduced with 22 g. IIA yielded (XIV), CGH9N50.0.5H2504.0.5H20. XIV (1 g.) in 100 cc. water at 60' treated with 1 g. Ac2, and the mixt. heated 15 min. at 60-70', allowed to stand overnight, and condd. to 25 cc. yielded 0.22 g. 2-mathylamino-4-hydroxy-6,7-dimethylpteridine (XV), fine light yellow needles from water, m. 277-81'. The filtrate from 1 g. XV and 0.57 g. BaCl2 added to 1 g. Bz2 in 25 cc. alc., and the mixt. refluxed 2 hrs. yielded 2-mathylamino-4-hydroxy-6,7-diphenylpteridine (XVI), decomp. 346-54'. XVI (0.175 g.), 10 cc. POCl3, and 0.7 g. PCl5 refluxed 2 hrs., the POCl3 distd. off, and the residue poured onto ice yielded chlorinated XVI (XVII). XVII (1 g.) and 20 cc. MeOH (satd. with NN3 at 0') heated in a sealed tube 16 hrs. at 155' yielded 2.4-dismino-6,7-diphenylpteridine. XIV (8.4 g.) with IV and V yielded 2.4-dismino-6,7-diphenylpteridine. XIV (8.4 g.) with IV and V yielded 1.7 g. sulfate. XIV (25 g.) dissolved in 2.5 l. water beated to 100' yielded 1.0 g. NIII dissolved in 300 cc. water with a min. of dil. NaOH at 60' yielded 5.1 g. 3-mathyl-2,6-diamino-5-nitroso-(4H)-pyrimidone (XVII). XVII (2 g.) in 75 cc. water at 40' with 1 g. Ac2 yielded 1.1 g. 3-mathyl-2,6-diamino-6,7-diphenyl-(pyrimidone, began to sublime at 350-60', did not m. below 37

L8 ANSWER 33 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

(V.T.) (1.e., the maximum number of dilns. necessary before a feach suspension loses all vibriostatic activity) of 4% of IV in the diet was 160-320, tested against an inoculum of 103 vibrios of Vibrio cholerae per al., for 24 hrs. Under the same conditions the V.T. of 0.2, 0.4, and 0.8% dietary levels of I and II was, resp., <80, <80, and 80, 320, 320-640, and 5120, and <80 for all levels of III. Feces from diets containing 0.4% pteridine plus 3.6% of IV showed a V.T. of 1280, 2560, and 320 for I, II, and III, resp. The percentage of dry weight of the compds. in the faces was 0.3 and 0.9% for I at the low and high diet level, 0.1 and 1.0% for II, and 0.3 and 1.5% for II. The pteridines were estimated fluorometrically. Although chloroamphenicol (V) was powerfully vibriostatic in vitro when added to normal mouse fecal suspensions [min. vibriostatic concentration 0.5 y/ml.), there was no vibriostatic activity seen in the feces of mice fed 1.0% of V in the diet. For I, II, III, and IV, resp., the in vitro activities were, in y/ml., 15.5, 2.0, > 500, and 500; and 7.8 and 3.9 for I- and II-phosphates, resp.

17 694514-86-8, Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)- (SCI) (CA INDEX NAME)

CN Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)- (SCI) (CA INDEX NAME)

L8 ANSVER 34 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1951:45059 CAPLUS
DOCUMENT NUMBER: 45:45059
ORIGINAL REFERENCE NO.: 45:7691h

ITILE: Effect of 2,4-diamino-5-(p-chlorophenoxy)-6methylpyrimidine and 2,4-diamino-6,7-diphenylpteridine
on a chloroguanide-resistant strain of Plasmodium
gallinaceum

AUTHOR(S): Greenberg, Joseph; Richeson, Edna H.
CORPORATE SOURCE: Natl. Inst. Health, Bethesda, MD
Proceedings of the Society for Experimental Biology
and Nedicine (1951), 77, 174-6
CODEN: PSEBAA, ISSN: 0037-9727
DOCUMENT TYPE: Journal
LANGUAGE: Unavailable
AB A chloroguanide-resistant strain of Pl. gallinaceum was cross-resistant to
the second compound but not to the first. The first compound and
chloroguanide were not synergistic in their antimalarial activity.

IT 10101-83-6, Pteridine, 2,4-diamino-6,7-diphenyl(effect on choroguanide-resistant Plasmodium pallinaceum)
RN 10181-93-6 CAPLUS
CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

La ANSWER 35 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:

ORIGINAL REFERENCE NO: 45:33453

AUTHOR(S):

CORPORATE SOURCE:

CORPORATE SOURCE:

Allen and Hanburys, Ltd., Vare, UK
Annals of Tropical Medicine & Parasitology (1950), 44,
273-80

CODEN: ATMPA2; ISSN: 0003-4983

DOCUMENT TYPE:

LANGUAGE:

LANGUAGE:

AUTHOR (S):

CORPORATE SOURCE:

Allen and Hanburys, Ltd., Vare, UK
Annals of Tropical Medicine & Parasitology (1950), 44,
273-80

CODEN: ATMPA2; ISSN: 0003-4983

DOCUMENT TYPE:

LANGUAGE:

LANGUAGE:

LANGUAGE:

LANGUAGE:

AUTHOR (S):

CORPORATE SOURCE:

Annals of Tropical Medicine & Parasitology (1950), 44,
273-80

CODEN: ATMPA2; ISSN: 0003-4983

DOCUMENT TYPE:

Journal LANGUAGE:

AND LANGUAGE:

LANGUAGE:

AND LANGUAGE:

LANGUAGE:

AND LANGUAGE:

AND LANGUAGE:

AND LANGUAGE:

AND LANGUAGE:

LANGUAGE:

AND LANGUAGE:

increasing PGA concus. in the case of pteridines II, III, IV, and V, but rising for I. Some antagonism towards II and IV was shown by PA at 0.02 to 2 \( \gamma \) with V. cholerae 100 \( \gamma \) mid of PGA overcame to some extent the inhibitory activity of the pteridines and of sulfaquanidine (VI), alone or in combination. Ten \( \gamma \) mid PGA had no effect. PA at 100 \( \gamma \) mil. also overcame the vibriostatic action of II and IV, but 1 \( \gamma \) mil. sign overcame the vibriostatic action of II and IV, but 1 \( \gamma \) mil. was ineffective. PABA (0.001 to 0.1 \( \gamma \) mil. did not antagonize the activity of the pteridines, but was effective against VI. Peptone, at 100 \( \gamma \) mil. readily antagonized the vibriostatic action of VI, and to a lesser extent that of II, but had no effect on action of IV. 18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl-(in cholera therapy) 18181-93-6 CAPLUS 2,4-Pteridinediamine, 6,7-diphenyl-(QCI) (CA INDEX NAME)

IT

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ &$$

L8 ANSWER 37 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 155:33451 CAPLUS
OCCUMENT NUMBER: 45:33451
ORIGINAL REFERENCE NO.: 45:58151,5816a-c
TITLE: Chemotherapy of cholera. II. In vitro vibriostatic properties of certain 2.4-diaminopteridines
CORPORATE SOURCE: CONFORATE SOURCE: Allen and Hamburya, Ltd., Ware, UK
Annals of Tropical Medicine & Parasitology (1950), 44, 156-60
CODEN: ATMPA2; ISSN: 0003-4983
Journal

DOCUMENT TYPE: LANGUAGE:

CODEN: ATMPA2; ISSN: 0003-4983

JURNT TYPE: Journal

JUAGE: Unavailable

of. C.A. 44, 6521d. 6,7-Disubstituted 2,4-diaminopteridines were prepared
and tested for vibriostatic activity. Of the 2,4-diamino-6,7dialkylpteridines the diisopropyl (I) and di-sec-Bu (II) compds. were the
most active but the di-Et and di-Pr derivs. were also active. Generally,
alkyl substituents of less than 2 or more than 3 C atoms were inactive.
In the 6,7-diaryl series di(p-methoxyphenyl) and di(1-furyl) derivs. were
active but the di(o-methoxyphenyl), di-Ph, and dibenzyl compds.

were not. With condensed ring substituents at the 6,7-positions, the most
effective was 2,4-diamino-1'-methylindolo-(2',3',6',7)pteridine (III). In
all other compds. tested the min. inhibitory concentration rose markedly as

incubation time increased. The 1'-Et and 1'-propylindolo-(2',3',6,7) compds. were active at about 20-40 y/mL. Sulfaquanidine (IV) was used for comparison. Only III remained fully as effective against 106 as against 103 vibrios/mL. All strains of vibrios tested were inhibited by III. There was no difference in activity of I in a synthetic medium as compared to a peptone broth, while the min. inhibitory concentration of IV

lower in the synthetic medium. The phosphate, Cl-, and NO3- salts of III were prepared and tested at various pH values in the synthetic medium. The phosphate showed good activity from pH 7 to 8.5, the growth range for vibrios. The solubilities in NEO at 37' and pH 7 of several of the compds. (in mg./mL.) were: I, 0.1r I-phosphate, 18.7; II, 0.12; II-phosphate, 27.7; III, 0.04 and III-phosphate, 0.58.
694514-86-8, Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)(in cholera therapy)
694514-86-8 CAPLUS
Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)- (SCI) (CA INDEX NAME)

L8 ANSVER 36 OF 45 CAPLUS COPYRIGHT 2006 ACS ON STN
ACCESSION NUMBER: 1951:33452 CAPLUS
ORIGINAL REFERENCE NO.: 45:33452
ORIGINAL REFERENCE NO.: 45:5916C-e
TITLE: Chemotherapy of cholera. III. The action of preridine-sulfonamide mixtures upon Vibrio cholerae and upon the mouse
AUTHOR(S): Collier, H. O. J.; Hall, Iris F.; Waterhouse, Pamela D.

CORPORATE SOURCE: SOURCE:

D. Allen and Hanburys, Ltd., Ware, UK Annals of Tropical Medicine & Parasitology (1950), 44, 161-7 CODEN: ATMPA2; ISSN: 0003-4983

CODEN: ATHPA2; ISSN: 0003-4983

DOCUMENT TYPE:

Journal

AB The 2,4-diamino-6,7-diethyl, dipropyl, diisopropyl (I), di-sec-butyl, di(1-f-uryl), and di(p-methoxyphenyl) (II) pteridines, 2,4-diaminocamphano-(2',3',6,7 or 7,6) pteridine, and 2,4-diamino-1'-methylindolo-(2',3',6,7)-pteridine (III) were tested with and without sulfaquanidine (IV). All showed marked spaergism with IV. A mixture composed of 10% pteridine and 90% IV had about the same activity as the pure pteridine for incubation periods up to 24 hrs. The mixts. were generally more active than the pure pteridines upon longer incubation. The LDSO (in mg./kg., intraperitoneally in mice) of 1, II, and III, resp., was: 141, 186, and 126, the LDSO of IV was 870. The LDSO of 10% mixts. of I, II, and III with IV were, resp., 1023, 890, and 878 mg./kg.

II 694514-86-8, Zteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)(in cholera therapy)

RN 694514-86-8 CAPLUS

Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)- (CA INDEX NAME)

L8 ANSWER 38 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1550:43971 CAPLUS
OCCUMENT NUMBER: 44:43971
ORIGINAL REFERENCE NO.: 44:8417b-c
ITILE: AUTHOR(5): Vibriostatic activity in certain series of pteridines
AUTHOR(5): Collier, H. O. J.; Campbell, N. R.; Fitzgerald, M. E.

H. Allen & Hanburys, Ltd., Ware, Herts, UX Nature (London, United Kingdom) (1950), 165, 1004-5 CODEN: NATUAS; ISSN: 0028-0836 CORPORATE SOURCE: SOURCE:

CODEN: NATUAS; ISSN: 0028-0836

DOCUMENT TYPE:
Journal
Journal
AB Condensations of tetraminopyrimidine with N-methylisatin in the presence
of mineral acid gives a mixture of an active vibriostatic isomer 2,
4-diamino-1'-methylimdole-(2', 3', 6, 7)-pteridine. In tests, (against
Vibrio cholerae) the activity was greatest in the disopropyl compound

IT 18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl(90514-86-8, Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)(vibriostatic activity of)
RN 18181-93-6 CAPLUS
CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

694514-86-8 CAPLUS
Pteridine, 2,4-diamino-6,7-bis(p-methoxyphenyl)- (5CI) (CA INDEX NAME)

L8 ANSWER 39 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1950:28053 CAPLUS
OCCUMENT NUMBER: 44:2806-3
TITLE: 44:54806-7
ANTHOR(S): 50:5053 CAPLUS
AUTHOR(S): 60:5053 CAPLUS
AUTHOR(S): 60:5053 CAPLUS
AUTHOR(S): 50:5053 CAPLUS
AUTHOR(S): 60:5053 CAPLUS
AUTHOR
AUTHOR(S): 60:5053 CAPLUS
AUTHOR
AUTHOR(S): 64:5053 CAPLUS
AUTHOR(S): 62:5053 CAPLUS
AUTHOR(S): 62:5053 CAPLUS
AUTHOR
AUTHOR(S): 62:5053 CAPLUS
AUTHOR(S): 62:5053 CAPLUS
AUTHOR
AUTHOR(S): 62:5053 CAPLUS
AUTHOR
AUT

doses of Sulfabizine. Data, as was much more toxic than compds.

18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl151648-52-1, Pteridine, 2,4-diamino-6,7-bis(p-aminophenyl)(antimalarial activity of)
18181-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

151648-52-1 CAPLUS 2,4-Pteridinediamine, 6,7-bis(4-aminophenyl)- (9CI) (CA INDEX NAME)

L8 ANSWER 40 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1950:14970 CAPLUS
DOCUMENT NUMBER: 44:14970
ORIGINAL REFERENCE NO.: 44:2992b-d
ITILE: A new synthesis of pteridines
TITLE: A new synthesis of pteridines
TITLE: A new synthesis of pteridines
OURCE: NATURAS: ISSN: 0028-0836
DOCUMENT TYPE: Journal
LANGUAGE: Unavailable
GI For diagram(s), see printed CA Issue.
AB Isomer formation and ambiguity about the structure of the product arising
from the condensation of 4,5-diaminopyrimidines with diketones or other
suitable compds. are avoided by using 5-nitroso-4-aminopyrimidines and
ketones as reactants. The following derives of I have been made by
condensation in HOAc at 100-60'. RI, R2 - NHZ, R3 R4 - Ph, m.
282'R1, R2 - NHZ, R3 - Ph, R4 - Me, m. 330', R1, R2 - PHZ,
R3 R4 - -CO.NH.CO.NH-, absorption maximum, \$264, 369 (log .vepsiln.
4.11, 4.34), min., \$294 (log .vepsiln. 3.59); and R1, R2 - OH,
R3R4 - -CO.NH.CO.NH-, absorption maximum, \$280, 388 (log .vepsiln.
4.2, 4.3), min., \$275, 322 (log .vepsiln. 4.16, 3.4).

IT 18191-93-6, Pteridine, 2,4-diamino-6,7-dipheny1(preparation of)
RN 18181-93-6 CAPLUS
CN 2,4-Pteridinediamine, 6,7-dipheny1- (9CI) (CA INDEX NAME)

ANSWER 39 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) 10101-93-6, Pteridine, 2,4-diamino-6,7-diphenyl-(antimalarial activity of, and its potentiation by sulfadiazine and inhibition by folic acid) 10101-93-6 CAPLUS 2,4-Pteridinediamina. 6 2-44

4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

L8 ANSWER 41 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1949:47440 CAPLUS
ORIGINAL REFERENCE NO.: 43:8755e-h
TITLE: 43:8555e-h
Hematologic effect in rate of pterins structurally related to pteroylglutamic acid
AUTHOR(S): Swendseid, Marian E., Wittle, E. L., Moersch, G. W., Bird, O. D., Brown, Raymond A.
SOURCE: Journal of Biological Chemistry (1949), 179, 1175-82
CODEN: JECHA3; ISSN: 0021-9258

CODEN: JECHA3; ISSN: 0021-9258

JOURNAL TYPE: JOURNAL JURGE JURG

of I/100 g. developed laucopenia with agranulocytomia, but there was no effect on hemoglobin concentration The leucopenia was prevented by the addition of

addition of
an equivalent amount of ptercylglutamic acid (VI). II and III, at 50 mg.

thad
no effect on hematologic pattern, but II at 500 mg. t gave results similar
to those with I. IV caused leucopenia with agranulocytosis and also
anemia when fed at a level of 0.3 mg. t. The changes were prevented by
the addition of an equivalent amount of VI. The effect of V was similar to

t of IV, but the dietary level required was much higher (500 mg. %). II and at lower levels, prevented the agranulocytosis caused by the feeding of sulfasuxidine. 18181-93-6. Pteridine, 2,4-diamino-6,7-diphenyl-(hematologic effect of) 18181-93-6 CAPLUS 2,4-Pteridinediamine, 6,7-diphenyl-(9CI) (CA INDEX NAME)

ΙŢ

L8 ANSWER 42 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1949:47037 CAPLUS
DOCUMENT NUMBER: 43:47037
ORIGINAL REFERENCE NO.: 43:8491c-e
TITLE: growth by folic acid antagonists
Hertz. Roy: Tullner, Vm. W.
AUTHOR(S): Hertz. Roy: Tullner, Vm. W.
SOURCE: Endocrisology (1949), 44, 278-82
CODEN: ENDOAD, 155N: 0013-7227
DOCUMENT TYPE: Journal
LANGUAGE: A. 42, 4659a. In stilbestrol-treated chicks and estradioltreated ovariectomized rate, quant. inhibition of estrogen-induced tissue growth in the female genital tract was obtained with the folic acid antagonists, 4-aninopteroylespartic, 4-desoxypteroylglutamic, and 4-anino-N10-methylpteroylglutamic acids, 2,4-diamino-6,7-dimethylpteridine, 2,4-diamino-6,7-dimethylpteridine, The inhibition was reversed by administration of folic acid.

IT 18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl-teridine (2,4-diamino-6,7-diphenylpteridine)
RN 18181-93-6 CAPLUS
CN 2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

ANSVER 43 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) retention of antifolic acid activity. The introduction into I of any of the solubilizing groups investigated results in some lowering of antifolic acid activity; the effect of certain structural changes in I on such activity is discussed.

18181-93-6, Pteridine, 2,4-diamino-6,7-diphenyl(and derivs.)

18181-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

6967-77-7. Pteridine, 2,4-diamino-6,7-bis(p-hydroxyphenyl)151648-32-1. Pteridine, 2,4-diamino-6,7-bis(p-sminophenyl)80455-05-3. Acstanilide, 4'-(7-(p-sectamidophenyl)-2,4-diamino-6pteridyl]- 855629-16-2. Phenol, p-[2,4-diamino-7-(phydroxyphenyl]-6-pteridyl]- 855868-32-9. Hethanol,
[p-[2,4-diamino-7-(p-[dhydroxynethyl)amino]phenyl]-6-pteridyl]anilino]857228-86-5. Pteridine, 2,4-diamino-6,7-bis[a-sminophenyl][preparation of]
6567-77-7 CAPLUS
Phenol, 4,4'-(2,4-diamino-6,7-pteridinediyl)bis- (9CI) (CA INDEX NAME)

151648-52-1 CAPLUS 2,4-Pteridinediamine, 6,7-bis(4-aminophenyl)- (9CI) (CA INDEX NAME)

804555-05-3 CAPLUS Acetanilide, 4'-[7-(p-acetanidophenyl)-2,4-diamino-6-pteridyl]- [5CI] (CA INDEX NAME)

Page 25 Saeed

L8 ANSVER 43 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1949:22617 CAPLUS
DOCUMENT NUMBER: 43:22617
ORIGINAL REFERENCE NO.: 43:4268=-1,4269=-c
TITLE: Pteridines. IV. Derivatives of 2,4-diamino-6,7-diphenylpteridine.
Cain. C. K.; Taylor, E. C., Jr.; Daniel, Louise J.
JOURNEL: Journal of the American Chemical Society (1949), 71, 892-6

COURN JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal
LANGUAGE: Unavailable
AB cf. C.A. 43, 654d. 2,4-Dlamino-6,7-diphenylpteridine (I) (1 g.) and 50 m.
L. Ac20, refluxed 16 h., give 55% of the N4-Ac derivative, light yellow, m. slowly at 140-50°, 0.5 g. I, 10 mL. Ac20, and 3 mL. H2504, heated 1
h. on the steam bath, give 68% of the N2,N4-di-Ac derivative, light yellow, decompose slowly above 190°. 2-Amino-4-bydromy-6,7-diphenylpteridine (II) (1 g.), 60 mL. PCC13, and 5 g. PC15, refluxed 2 h., give 81% of the 4-C1 compound (III), bright yellow, could not be crystallized because of hydrolysis to II. III (1 g.), 10 mL. MeNH2, and 30 mL. ECOH, heated 16 h. at 155°, give 27% 2-amino-4-methylamino-6,7-diphenylpteridine, bright yellow, m. 237-8° (corrected), 2,4,5,6-cm sulfate (2 g.) in 90 mL. H200, treated with 2 g. [p-H2NCGH4CO)2,H2504 and refluxed 1 h., gives 1.73 p. 2,4-diamino-6,7-bis(p-aminophenyl)pteridine (IV), bright orange, decompose 308-5° (corrected), 2,4,5,6-cm serialized branches (IV), bright orange, decompose 308-5° (corrected), 2,4,5,6-cm serialized branches (IV), m. 234-7' (corrected), IV (0.1 g.) in 4 mL. bobling H20 containing sufficient concentrated HC1 to cause solution, treated with 0.2 and adjusted to pH 7.5 with NaHCO3, gives a quant. yield of 2,4-diamino-6,7-bis(p-((hydroxymethyl)) anino)phenyl)pteridine (VI). does

CONTENTS of the Content of the Conte

min.

Tal. H20 and 0.8 ml. concentrated H2DUE, treated with miles of and 1 h. at 100°, gives 71% 2,4-diamino-6,7-bis(p-hydroxyphenyllperidine, yellow; it was prepared also from (p-HOCGH4CO)2 and the bisulfite (ViII) of V (84%). V (3 9.) and 2 9. (m-O2NCGH4CO)2 in 70 ml. EtOH and 15 ml. ACET, refluxed 3 h., give a quant, yield of 2,4-diamino-6,7-bis(m-nitrophenyl)pteridine, m. 307-8' (corrected); catalytic reduction gives 65% of the m-aminophenyl compound, orange-yellow, decompose above 180°. The m-isomer of VII (85% yield) is hygroscopic and rapidly forms a trihydrate in the air. VIII (5 9.) in 20 ml. 0.5% NAOH, added to 5 9, phenanthrenequinone-3-sulfonic acid in 130 ml. H2O and refluxed 30 min., gives 88% 2,4-diaminophenanthro[9,10-e]pteridine-8 (or 11)-sulfonic acid, light yellow; does not m. up to 360°. The absorption spectra of these compds., qual. solubility in H2O, EtOH, and 0.1

HCl and NaOH, and their inhibitory indexes against Streptococcus faecalis are given. A (sulfinomethyl) amino group confers H2O solubility with the

ANSWER 43 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

855629-16-2 CAPLUS
Phenol, p-[2,4-diamino-7-[o-hydroxyphenyl]-6-pteridyl]- (SCI) (CA INDEX NAME)

855868-52-9 CAPLUS
Methanol, [p-[2,4-diamino-7-[p-[(hydroxymethyl)amino]phenyl]-6pteridyl]anilino]- (SCI) (CA INDEX NAME)

857228-86-5 CAPLUS
Pteridine, 2,4-diamino-6,7-bis[m-aminophenyl]- (5CI) (CA INDEX NAME)

ANSWER 43 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

857398-11-9 CAPLUS Pteridine, 2,4-diamino-6,7-bis[m-nitrophenyl]- (SCI) (CA INDEX NAME)

L8 ANSWER 45 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1947:31103 CAPLUS DOCUMENT NUMBER: 41:31103 ORIGINAL REFERENCE NO.: 41:6258a-f

ORIGINAL REFERENCE NO.: 4116289s-f
TITLE: Pyrimido(4,5-b)pyrazines. II. 2,4-Diaminopyrimido(4,5-b)pyrazine and derivatives
b)pyrazine and derivatives
b)pyrazine and derivatives
corporate source: Corporate sourc

treated with 1 g. accmaphthenequinone in 25 ml. HCONMe2 and the mixture heated 4 hrs. on the steam bath, give 90, \$ 2,4-disminoaccmaphtho[1,2-elpyrimido[4,5-b]pyrazine, needles, decompose on heating. I (2 g.), 1.5 g. phenanthreneqvinone, 250 ml. 95% EtOH, and 5 ml. 10% aqueous NaOH, refluxed

hrs., give 84% 2,4-diaminophenanthro[9,10-e]pyrimido[4,5-b]pyrazine, needles, sinters 340° without melting. I (15 g.), 6 g. AcCHD, and 200 cc. H2O give 90% of the 6(or 7)-He derivative of II, prisms, decompose

heating. All these compds. show parallel extinction. The ultraviolet absorption spectra are given of the above compds. and of some reported in Part I. Several derivs. of II exhibit marked antifolic acid activity for several bacteria.

1810-93-6, Pteridine, 2,4-diamino-6,7-diphenyl-(preparation of) 18181-93-6 CAPLUS
2,4-Pteridinediamine, 6,7-diphenyl- (9CI) (CA INDEX NAME)

L8 ANSWER 44 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 1947:37525 CAPLUS
COCUMENT NUMBER: 41:7525 CAPLUS
CRIGINAL REFERENCE NO.: 41:74381,74393-b
TITLE: Studies with Streptococcus faecalis, Lactobacillus casei, and Lactobacillus arabinosus
AUTHOR(S): Daniel, Louise J.: Norris, L. C.: Scott, M. L.:
Heuser, G. F.
CORPORATE SOURCE: Cornell Univ., Ithaca
SOURCE: JOURNAL of Source Journal of Biological Chemistry (1947), 169, 689-97
CODEN: JECHAJ; ISSN: 0021-9258

DOCUMENT TYPE: Journal
LANGUAGE: Unavailable
AB The following synthetic pterins were used: 2,4-diamino-6,7dimethylpyrimido-(4,5-b)pyrazine, 2,4-diamino-6,7dimethylpyrimido-(4,5-b)pyrazine, 2,4-diamino-6,7diphenylpyrimido(4,5-b)pyrazine, 2,4-diamino-6,7diphenylpyrimido(4,5-b)pyrazine, 2,4-diamino-6,7diaminopananthro(9,10-e)pyrimido(4,5-b)pyrazine, 2,4-diamino-6,7diaminopananthro(9,10-e)pyrimido(4,5-b)pyrazine, 2,4-diaminosus, which synthesizes its own I. The secalis and L. casei which
require folic acid (1) as an essential nutrient, but also for L.
arabinosus, which synthesizes its own I. The substitution of OH for NHZ
in the 4- or 2-position destroyed the anti-1 activity. Those pterins with
4-NHZ groups varied in anti-1 with the nature of the substitution in the
6- and 7-positions.

1 18181-93-6 CAPLUS

N1 18181-93-6 CAPLUS

CN 2,4-Pteridine, 2,4-diamino-6,7-diphenyl(9COL) (CA INDEX NAME)

L8 ANSWER 45 OF 45 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

=> logoff

ALL L# QUERIES AND ANSWER SETS ARE DELETED AT LOGOFF

LOGOFF? (Y)/N/HOLD:Y

SINCE FILE TOTAL ENTRY SESSION 230.87 565.88 COST IN U.S. DOLLARS

FULL ESTIMATED COST

TOTAL DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE

ENTRY SESSION -33.75 CA SUBSCRIBER PRICE

STN INTERNATIONAL LOGOFF AT 08:48:39 ON 30 MAY 2006